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(57) Abstract

Gene sequences as shown in SEQ ID NOS:1-85 have been found to be significantly associated with metastatic potential of cancer cells, especially breast and colon cancer cells. Methods are provided for determining the risk of metastasis of a tumor, which involve determining whether a tissue sample from a tumor expresses a polypeptide encoded by a gene as shown in SEQ ID NOS:1-85, or a substantial portion thereof.

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METASTATIC BREAST AND COLON CANCER REGULATED GENES

TECHNICAL FIELD OF THE INVENTION

This invention relates to methods for predicting the behavior of tumors. More particularly, the invention relates to methods in which a tumor sample is examined for expression of a specified gene sequence thereby to indicate propensity for metastatic spread.

BACKGROUND OF THE INVENTION

Breast cancer is one of the most common malignant diseases in women, with about 1,000,000 new cases per year worldwide. Colon cancer is another of the most common cancers. Despite use of a number of histochemical, genetic, and immunological markers, clinicians still have a difficult time predicting which tumors will metastasize to other organs. Some patients are in need of adjuvant therapy to prevent recurrence and metastasis and others are not. However, distinguishing between these subpopulations of patients is not straightforward, and course of treatment is not easily charted. There is a need in the art for new markers for distinguishing between tumors which will or have metastasized and those which are less likely to metastasize

SUMMARY OF THE INVENTION

It is an object of the present invention to provide markers for distinguishing between tumors which will or have metastasized and those which are less likely to metastasize. These and other objects of the invention are provided by one or more of the embodiments described below.

One embodiment of the invention provides an isolated and purified human protein having an amino acid sequence which is at least 85% identical to an amino acid sequence encoded by a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.

Another embodiment of the invention provides a fusion protein which comprises a first protein segment and a second protein segment fused to each other by

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means of a peptide bond. The first protein segment consists of at least six contiguous amino acids selected from an amino acid sequence encoded by a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.

Yet another embodiment of the invention provides an isolated and purified polypeptide consisting of at least six contiguous amino acids of a human protein having an amino acid sequence encoded by a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.

Still another embodiment of the invention provides a preparation of antibodies which specifically bind to a human protein which comprises an amino acid sequence encoded by a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.

Even another embodiment of the invention provides an isolated and purified subgenomic polynucleotide comprising at least 11 contiguous nucleotides of a nucleotide sequence which is at least 96% identical to a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.

Another embodiment of the invention provides an isolated and purified gene which comprises a coding sequence comprising a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.

Yet another embodiment of the invention provides a method for determining metastasis in a tissue sample. An expression product of a gene which comprises a coding sequence selected from the group consisting of SEQ ID NOS:1, 2, 4, 5, 9, 11, 13, 14, 18, 19, 20, 22, 24, 26, 29, 30, 33, 35, 36, 38-41, 45, 48, 52, 55, 57, 58, 60, 63-66, 69-74, 76, 80, 82, and 83 is measured in a tissue sample. A tissue sample which expresses the product is categorized as metastatic.

Still another embodiment of the invention provides a method for determining metastasis in a tissue sample. An expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:3, 7. 8, 10. 12, 15-17, 21, 23, 28, 31, 34, 37, 42-44, 46, 47, 49-51, 53, 59, 61, 62, 67, 68, 75, 77-79, 81, 84, and 85 is measured in a tissue sample. A tissue sample which does not express the product is categorized as metastatic.

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Even another embodiment of the invention provides a method for determining metastatic potential in a tissue sample. An expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:1, 2, 4, 5, 9, 11, 13, 14, 18, 19, 20, 22, 24, 26, 29, 30, 33, 35, 36, 38-41, 45, 48, 52, 55, 57, 58, 60, 63-66, 69-74, 76, 80, 82, and 83 is measured in a tissue sample. A tissue sample which expresses the product is categorized as having metastatic potential.

A further embodiment of the invention provides a method for determining metastatic potential in a tissue sample. An expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:3, 7, 8, 10, 12, 15-17, 21, 23, 28, 31, 34, 37, 42-44, 46, 47, 49-51, 53, 59, 61, 62, 67, 68, 75, 77-79, 81, 84, and 85 is measured in a tissue sample. A tissue sample which does not express the product is categorized as having metastatic potential.

Another embodiment of the invention provides a method of predicting the propensity for metastatic spread of a breast tumor preferentially to bone or lung. An expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NO:1, 5, 11, 18, 20, 22, 24, 30, 33, 35, 36, 38, 45, 52, 58, 65, 66, 70, 74, 76, and 80 is measured in a breast tumor sample. A breast tumor sample which expresses the product is categorized as having a propensity to metastasize to bone or lung.

Even another embodiment of the invention provides a method of predicting propensity for metastatic spread of a breast tumor preferentially to lung. An expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:2, 4, 9, 13, 14, 19, 26, 29, 39-41, 48, 55, 57, 60, 63, 64, 72, 73, 82, and 83 is measured in a breast tumor sample. A breast tumor sample which expresses the product is characterized as having a propensity to metastasize to lung.

Still another embodiment of the invention provides a method of predicting propensity for metastatic spread of a colon tumor. An expression product of a gene which comprises the nucleotide sequence shown in SEQ ID NO:56 is measured in a colon tumor sample. A colon tumor sample which expresses the product is characterized as having a low propensity to metastasize.

Even another embodiment of the invention provides a method for determining metastasis in a tissue sample. An expression product of a gene which comprises a coding sequence selected from the group consisting of SEQ ID NOS:3, 7, 8, 10, 12, 15-17, 21, 23, 25, 28, 31, 34, 37, 42-44, 46, 47, 49, 61, 62, 67, 68, 75, 77-79, 81, 84, and 85 is measured in a tissue sample. A tissue sample which expresses the product is categorized as non-metastatic.

Yet another embodiment of the invention provides a method for determining metastasis in a tissue sample. An expression product of a gene which comprises a coding sequence selected from the group consisting of SEQ ID NOS:3, 7, 8, 10, 12, 15-17, 21, 23, 25, 28, 31, 34, 37, 42-44, 46, 47, 49, 61, 62, 67, 68, 75, 77-79, 81, 84, and 85 is measured in a tissue sample. A tissue sample which does not express the product is categorized as metastatic.

The invention thus provides the art with a number of genes and proteins, which can be used as markers of metastasis. These are useful for more rationally prescribing the course of therapy for breast or colon cancer patients.

DETAILED DESCRIPTION

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It is a discovery of the present invention that a number of genes are differentially expressed between metastatic cancer cells, especially cancer cells of the breast and colon, and non-metastatic cancer cells. These genes are metastatic marker genes. This information can be utilized to make diagnostic reagents specific for the expression products of the differentially expressed genes. It can also be used in diagnostic and prognostic methods which will help clinicians in planning appropriate treatment regimes for cancers, especially of the breast or colon.

Some of the polynucleotides disclosed herein represent novel genes which are differentially expressed between non-metastatic cancer cells and cancer cells which have a potential to metastasize. SEQ ID NOS:1-63 represent novel metastatic marker genes (Table 1). SEQ ID NOS:64-85 represent known genes which have been found to be differentially expressed in metastatic relative to non-metastatic cancer cells (Table 2). Some of the metastatic marker genes disclosed herein are expressed in

metastatic cells relative to non-metastatic cells, particularly in breast cancer cells which metastasize to bone and lung (SEQ ID NOS:1, 5, 11, 18, 20, 22, 24, 30, 33, 35, 36, 38, 45, 52, 58, 65, 66, 70, 74, 76, and 80). One metastatic marker gene (SEQ ID NO:56) is expressed in non-metastatic breast cancer cells and in colon cancer cells with low metastatic potential. Other metastatic marker genes are expressed in metastatic cancer cells, particularly in breast cancer cells which metastasize only to lung (SEQ ID NOS:2, 4, 9, 13, 14, 19, 26, 29, 39-41, 48, 55, 57, 60, 63, 64, 72, 73, 82, and 83). Still other metastatic marker genes (SEQ ID NOS:3, 7, 8, 10, 12, 15-17, 21, 23, 28, 31, 34, 37, 42-44, 46, 47, 49, 61, 62, 67, 68, 75, 77-79, 81, 84, and 85) are expressed in cancer cells which do not typically metastasize, particularly in breast cancer cells. Identification of these relationships and markers permits the formulation of reagents and methods as further described below. Other metastatic marker genes, such as those which comprise a nucleotide sequence shown in SEQ ID NOS:6, 27, 32, and 54, can be used to identify cancerous tissue, particularly breast cancer tissue.

Sequences of metastatic marker genes are disclosed in SEQ ID NOS:1-85. Metastatic marker proteins can be made by expression of the disclosed polynucleotide molecules. Amino acid sequences encoded by novel polynucleotides of the invention can be predicted by running a translation program for each of three reading frames for a disclosed sequence and its complement. Complete polynucleotide sequences can be obtained by chromosome walking, screening of libraries for overlapping clones, 5' RACE, or other techniques well known in the art.

Reference to metastatic marker nucleotide or amino acid sequences includes variants which have similar expression patterns in metastatic relative to non-metastatic cells, as described below. Metastatic marker polypeptides can differ in length from full-length metastatic marker proteins and contain at least 6, 8, 10, 12, 15, 18, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 120, 140, 160, 180, or 200 or more contiguous amino acids of a metastatic marker protein.

Variants of marker proteins and polypeptides can also occur. Metastatic marker protein or polypeptide variants can be naturally or non-naturally occurring. Naturally occurring metastatic marker protein or polypeptide variants are found in

humans or other species and comprise amino acid sequences which are substantially identical to the proteins encoded by genes corresponding to the nucleotide sequences shown in SEQ ID NOS:1-85 or their complements. Non-naturally occurring metastatic marker protein or polypeptide variants which retain substantially the same differential expression patterns in metastatic relative to non-metastatic cancer cells as naturally occurring metastatic marker protein or polypeptide variants are also included here. Preferably, naturally or non-naturally occurring metastatic marker protein or polypeptide variants have amino acid sequences which are at least 85%, 90%, or 95% identical to amino acid sequences encoded by the nucleotide sequences shown in SEQ ID NOS:1-85. More preferably, the molecules are at least 98% or 99% identical. Percent sequence identity between a wild-type protein or polypeptide and a variant is determined by aligning the wild-type protein or polypeptide with the variant to obtain the greatest number of amino acid matches, as is known in the art, counting the number of amino acid matches between the wild-type and the variant, and dividing the total number of matches by the total number of amino acid residues of the wild-type sequence.

Preferably, amino acid changes in metastatic marker protein or polypeptide variants are conservative amino acid changes, *i.e.*, substitutions of similarly charged or uncharged amino acids. A conservative amino acid change involves substitution of one of a family of amino acids which are related in their side chains. Naturally occurring amino acids are generally divided into four families: acidic (aspartate, glutamate), basic (lysine, arginine, histidine), non-polar (alanine, valine, leucine, isoleucine, proline, phenylalanine, methionine, tryptophan), and uncharged polar (glycine, asparagine, glutamine, cystine, serine, threonine, tyrosine) amino acids. Phenylalanine, tryptophan, and tyrosine are sometimes classified jointly as aromatic amino acids.

It is reasonable to expect that an isolated replacement of a leucine with an isoleucine or valine, an aspartate with a glutamate, a threonine with a serine, or a similar replacement of an amino acid with a structurally related amino acid will not have a major effect on the biological properties of the resulting metastatic marker protein or polypeptide variant. Properties and functions of metastatic marker protein or polypeptide variants are of the same type as a metastatic marker protein or polypeptide comprising amino acid sequences encoded by the nucleotide sequences shown in SEQ ID NOS:1-85, although the properties and functions of variants can differ in degree. Whether an amino acid change results in a metastatic marker protein or polypeptide variant with the appropriate differential expression pattern can readily be determined. For example, nucleotide probes can be selected from the marker gene sequences disclosed herein and used to detect marker gene mRNA in Northern blots or in tissue sections, as is known in the art. Alternatively, antibodies which specifically bind to protein products of metastatic marker genes can be used to detect expression of metastatic marker proteins.

Metastatic marker variants include glycosylated forms, aggregative conjugates with other molecules, and covalent conjugates with unrelated chemical moieties. Metastatic marker variants also include allelic variants, species variants, and muteins. Truncations or deletions of regions which do not affect the differential expression of metastatic marker genes are also metastatic marker variants. Covalent variants can be prepared by linking functionalities to groups which are found in the amino acid chain or at the N- or C-terminal residue, as is known in the art.

Full-length metastatic marker proteins can be extracted, using standard biochemical methods, from metastatic marker protein-producing human cells, such as metastatic breast or colon cancer cells. An isolated and purified metastatic marker protein or polypeptide is separated from other compounds which normally associate with a metastatic marker protein or polypeptide in a cell, such as certain proteins, carbohydrates, lipids, or subcellular organelles. A preparation of isolated and purified metastatic marker proteins or polypeptides is at least 80% pure; preferably, the preparations are 90%, 95%, or 99% pure.

Metastatic marker proteins and polypeptides can also be produced by recombinant DNA methods or by synthetic chemical methods. For production of recombinant metastatic marker proteins or polypeptides, coding sequences selected from the nucleotide sequences shown in SEQ ID NOS:1-85, or variants of those

sequences which encode metastatic marker proteins, can be expressed in known prokaryotic or eukaryotic expression systems (see below). Bacterial, yeast, insect, or mammalian expression systems can be used, as is known in the art.

Alternatively, synthetic chemical methods, such as solid phase peptide synthesis, can be used to synthesize a metastatic marker protein or polypeptide. General means for the production of peptides, analogs or derivatives are outlined in Chemistry and Biochemistry of Amino Acids. Peptides, and Proteins -- A Survey of Recent Developments, Weinstein. B. ed., Marcell Dekker, Inc., publ., New York (1983). Moreover, substitution of D-amino acids for the normal L-stereoisomer can be carried out to increase the half-life of the molecule. Metastatic marker variants can be similarly produced.

Non-naturally occurring fusion proteins comprising at least 6, 8, 10, 12, 15, 18, 20, 25, 30, 35, 40, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 120, 140, 160, 180, or 200 or more contiguous metastatic marker amino acids can also be constructed. Human metastatic marker fusion proteins are useful for generating antibodies against metastatic marker amino acid sequences and for use in various assay systems. For example, metastatic marker fusion proteins can be used to identify proteins which interact with metastatic marker proteins and influence their functions. Physical methods, such as protein affinity chromatography, or library-based assays for protein-protein interactions, such as the yeast two-hybrid or phage display systems, can also be used for this purpose. Such methods are well known in the art and can also be used as drug screens.

A metastatic marker fusion protein comprises two protein segments fused together by means of a peptide bond. The first protein segment comprises at least 6, 8, 10, 12, 15, 18, 20, 25, 30, 35, 40, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 100, 120, 140, 160, 180, or 200 or more contiguous amino acids of a metastatic marker protein. The amino acids can be selected from the amino acid sequences encoded by the nucleotide sequences shown in SEQ ID NOS:1-85 or from variants of those sequences, such as those described above. The first protein segment can also comprise a full-length metastatic marker protein.

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The second protein segment can be a full-length protein or a protein fragment or polypeptide. The fusion protein can be labeled with a detectable marker, as is known in the art, such as a radioactive, fluorescent, chemiluminescent, or biotinylated marker. The second protein segment can be an enzyme which will generate a detectable product, such as β -galactosidase. The first protein segment can be N-terminal or C-terminal, as is convenient.

Techniques for making fusion proteins, either recombinantly or by covalently linking two protein segments, are also well known. Recombinant DNA methods can be used to prepare metastatic marker fusion proteins, for example, by making a DNA construct which comprises coding sequences selected from SEQ ID NOS:1-85 in proper reading frame with nucleotides encoding the second protein segment and expressing the DNA construct in a host cell, as described below.

Isolated and purified metastatic marker proteins, polypeptides, variants, or fusion proteins can be used as immunogens, to obtain preparations of antibodies which specifically bind to a metastatic marker protein. The antibodies can be used, *inter alia*, to detect wild-type metastatic marker proteins in human tissue and fractions thereof. The antibodies can also be used to detect the presence of mutations in metastatic marker genes which result in under- or over-expression of a metastatic marker protein or in expression of a metastatic marker protein with altered size or electrophoretic mobility.

Preparations of polyclonal or monoclonal antibodies can be made using standard methods. Single-chain antibodies can also be prepared. Single-chain antibodies which specifically bind to metastatic marker proteins, polypeptides, variants, or fusion proteins can be isolated, for example, from single-chain immunoglobulin display libraries, as is known in the art. The library is "panned" against metastatic marker protein amino acid sequences, and a number of single chain antibodies which bind with high-affinity to different epitopes of metastatic marker proteins can be isolated. Hayashi *et al.*, 1995, *Gene 160*:129-30. Single-chain antibodies can also be constructed using a DNA amplification method, such as the polymerase chain reaction

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(PCR), using hybridoma cDNA as a template. Thirion et al., 1996, Eur. J. Cancer Prev. 5:507-11.

Single-chain antibodies can be mono- or bispecific, and can be bivalent or tetravalent. Construction of tetravalent, bispecific single-chain antibodies is taught in Coloma and Morrison, 1997, *Nat. Biotechnol. 15*:159-63. Construction of bivalent, bispecific single-chain antibodies is taught in Mallender and Voss, 1994, *J. Biol. Chem.* 269:199-206.

A nucleotide sequence encoding the single-chain antibody can be constructed using manual or automated nucleotide synthesis, cloned into DNA expression constructs using standard recombinant DNA methods, and introduced into cells which express the coding sequence. as described below. Alternatively, single-chain antibodies can be produced directly using, for example, filamentous phage technology. Verhaar et al., 1995, Int. J. Cancer 61:497-501; Nicholls et al., 1993, J. Immunol. Meth. 165:81-91.

Metastatic marker-specific antibodies specifically bind to epitopes present in a full-length metastatic marker protein having an amino acid sequence encoded by a nucleotide sequence shown in SEQ ID NOS:1-85, to metastatic marker polypeptides, or to metastatic marker variants, either alone or as part of a fusion protein. Preferably, metastatic marker epitopes are not present in other human proteins. Typically, at least 6, 8, 10, or 12 contiguous amino acids are required to form an epitope. However, epitopes which involve non-contiguous amino acids may require more, e.g., at least 15, 25, or 50 amino acids.

Antibodies which specifically bind to metastatic marker proteins, polypeptides, fusion proteins, or variants provide a detection signal at least 5-, 10-, or 20-fold higher than a detection signal provided with other proteins when used in Western blots or other immunochemical assays. Preferably, antibodies which specifically bind to metastatic marker epitopes do not detect other proteins in immunochemical assays and can immunoprecipitate a metastatic marker protein, polypeptide, fusion protein, or variant from solution.

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Antibodies can be purified by methods well known in the art. Preferably, the antibodies are affinity purified, by passing the antibodies over a column to which a metastatic marker protein, polypeptide, variant, or fusion protein is bound. The bound antibodies can then be eluted from the column, for example, using a buffer with a high salt concentration.

Subgenomic polynucleotides contain less than a whole chromosome. Preferably, the polynucleotides are intron-free. In a preferred embodiment, the polynucleotide molecules comprise a contiguous sequence of 10, 11, 12, 15, 20, 25, 30. 32, 35, 40, 45, 50, 60, 70, 74, 80, 90, 100, 125, 150, 154, 175, 182, 200, 243, or 268 nucleotides selected from SEQ ID NOS:1-85 or the complements thereof. The complement of a nucleotide sequence shown in SEQ ID NOS:1-85 is a contiguous nucleotide sequence which forms Watson-Crick base pairs with a contiguous nucleotide sequence shown in SEQ ID NOS:1-85. The complement of a nucleotide sequence shown in SEQ ID NOS:1-85 (the antisense strand) is also a subgenomic polynucleotide, and can be used provide marker protein antisense oligonucleotides. Double-stranded polynucleotides which comprise one of the nucleotide sequences shown in SEQ ID NOS:1-85 are also subgenomic polynucleotides. Metastatic marker protein subgenomic polynucleotides also include polynucleotides which encode metastatic marker protein-specific single-chain antibodies and ribozymes, or fusion proteins comprising metastatic marker protein amino acid sequences.

Degenerate nucleotide sequences encoding amino acid sequences of metastatic marker protein and or variants, as well as homologous nucleotide sequences which are at least 85%, 90%, 95%, 98%, or 99% identical to the nucleotide sequences shown in SEQ ID NOS:1-85, are also metastatic marker subgenomic polynucleotides. Typically, homologous metastatic marker subgenomic polynucleotide sequences can be confirmed by hybridization under stringent conditions, as is known in the art. Percent sequence identity between wild-type and homologous variant sequences is determined by aligning the wild-type polynucleotide with the variant to obtain the greatest number of nucleotide matches, as is known in the art, counting the number of nucleotide matches between the wild-type and the variant, and dividing the total number of

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matches by the total number of nucleotides of the wild-type sequence. A preferred algorithm for calculating percent identity is the Smith-Waterman homology search algorithm as implemented in MPSRCH program (Oxford Molecular) using an affine gap search with the following search parameters: gap open penalty of 10, and gap extension penalty of 1.

Metastatic marker subgenomic polynucleotides can be isolated and purified free from other nucleotide sequences using standard nucleic acid purification techniques. For example, restriction enzymes and probes can be used to isolate polynucleotide fragments which comprise nucleotide sequences encoding a metastatic marker protein. Isolated and purified subgenomic polynucleotides are in preparations which are free or at least 90% free of other molecules.

Complementary DNA molecules which encode metastatic marker proteins can be made using reverse transcriptase, with metastatic marker mRNA as a template. The polymerase chain reaction (PCR) or other amplification techniques can be used to obtain metastatic marker subgenomic polynucleotides, using either human genomic DNA or cDNA as a template, as is known in the art. Alternatively, synthetic chemistry techniques can be used to synthesize metastatic marker subgenomic polynucleotides which comprise coding sequences for regions of metastatic marker proteins, single-chain antibodies, or ribozymes, or which comprise antisense oligonucleotides. The degeneracy of the genetic code allows alternate nucleotide sequences to be synthesized which will encode a metastatic marker protein comprising amino acid sequences encoded by the nucleotide sequences shown in SEQ ID NOS:1-85.

Purified and isolated metastatic marker subgenomic polynucleotides can be used as primers to obtain additional copies of the polynucleotides or as probes for identifying wild-type and mutant metastatic marker protein coding sequences. Metastatic marker subgenomic polynucleotides can be used to express metastatic marker mRNA, protein, polypeptides, or fusion proteins and to generate metastatic marker antisense oligonucleotides and ribozymes.

A metastatic marker subgenomic polynucleotide comprising metastatic marker protein coding sequences can be used in an expression construct. Preferably, the metastatic marker subgenomic polynucleotide is inserted into an expression plasmid (for example, the Ecdyson system, pIND, In Vitro Gene). Metastatic marker subgenomic polynucleotides can be propagated in vectors and cell lines using techniques well known in the art. Metastatic marker subgenomic polynucleotides can be on linear or circular molecules. They can be on autonomously replicating molecules or on molecules without replication sequences. They can be regulated by their own or by other regulatory sequences, as are known in the art.

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A host cell comprising a metastatic marker expression construct can then be used to express all or a portion of a metastatic marker protein. Host cells comprising metastatic marker expression constructs can be prokaryotic or eukaryotic. A variety of host cells are available for use in bacterial, yeast, insect, and human expression systems and can be used to express or to propagate metastatic marker expression constructs (see below). Expression constructs can be introduced into host cells using any technique known in the art. These techniques include transferrin-polycation-mediated DNA transfer, transfection with naked or encapsulated nucleic acids, liposome-mediated cellular fusion, intracellular transportation of DNA-coated latex beads, protoplast fusion, viral infection, electroporation, and calcium phosphate-mediated transfection.

A metastatic marker expression construct comprises a promoter which is functional in a chosen host cell. The skilled artisan can readily select an appropriate promoter from the large number of cell type-specific promoters known and used in the art. The expression construct can also contain a transcription terminator which is functional in the host cell. The expression construct comprises a polynucleotide segment which encodes all or a portion of the metastatic marker protein, variant, fusion protein, antibody, or ribozyme. The polynucleotide segment is located downstream from the promoter. Transcription of the polynucleotide segment initiates at the promoter. The expression construct can be linear or circular and can contain sequences, if desired, for autonomous replication.

Bacterial systems for expressing metastatic marker expression constructs include those described in Chang et al., Nature (1978) 275: 615, Goeddel et al., Nature (1979) 281: 544, Goeddel et al., Nucleic Acids Res. (1980) 8: 4057, EP 36,776, U.S. 4,551,433, deBoer et al., Proc. Nat'l Acad. Sci. USA (1983) 80: 21-25, and Siebenlist et al., Cell (1980) 20: 269.

Expression systems in yeast include those described in Hinnen et al., Proc. Nat'l Acad. Sci. USA (1978) 75: 1929; Ito et al., J. Bacteriol. (1983) 153: 163; Kurtz et al., Mol. Cell. Biol. (1986) 6: 142; Kunze et al., J. Basic Microbiol. (1985) 25: 141; Gleeson et al., J. Gen. Microbiol. (1986) 132: 3459, Roggenkamp et al., Mol. Gen. Genet. (1986) 202:302) Das et al., J. Bacteriol. (1984) 158: 1165; De Louvencourt et al., J. Bacteriol. (1983) 154: 737, Van den Berg et al., Bio/Technology (1990) 8: 135; Kunze et al., J. Basic Microbiol. (1985) 25: 141; Cregg et al., Mol. Cell. Biol. (1985) 5: 3376, U.S. 4,837,148, US 4,929,555; Beach and Nurse, Nature (1981) 300: 706; Davidow et al., Curr. Genet. (1985) 10: 380, Gaillardin et al., Curr Genet. (1985) 10: 49, Ballance et al., Biochem. Biophys. Res. Commun. (1983) 112: 284-289; Tilburn et al., Gene (1983) 26: 205-221, Yelton et al., Proc. Nat'l Acad. Sci. USA (1984) 81: 1470-1474, Kelly and Hynes, EMBO J. (1985) 4: 475479; EP 244,234, and WO 91/00357.

Expression of metastatic marker expression constructs in insects can be carried out as described in U.S. 4,745,051, Friesen et al. (1986) "The Regulation of Baculovirus Gene Expression" in: The Molecular Biology of Baculoviruses (W. Doerfler, ed.), EP 127,839, EP 155,476, and Vlak et al., J. Gen. Virol. (1988) 69: 765-776, Miller et al., Ann. Rev. Microbiol. (1988) 42: 177, Carbonell et al., Gene (1988) 73: 409, Maeda et al., Nature (1985) 315: 592-594, Lebacq-Verheyden et al., Mol. Cell. Biol. (1988) 8: 3129; Smith et al., Proc. Nat'l Acad. Sci. USA (1985) 82: 8404. Miyajima et al., Gene (1987) 58: 273; and Martin et al., DNA (1988) 7:99. Numerous baculoviral strains and variants and corresponding permissive insect host cells from hosts are described in Luckow et al., Bio/Technology (1988) 6: 47-55, Miller et al., in GENETIC ENGINEERING (Setlow, J.K. et al. eds.). Vol. 8 (Plenum Publishing, 1986), pp. 277-279, and Maeda et al., Nature, (1985) 315: 592-594.

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Mammalian expression of metastatic marker expression constructs can be achieved as described in Dijkema *et al.*, *EMBO J.* (1985) 4: 761, Gorman *et al.*, *Proc. Nat'l Acad. Sci. USA* (1982b) 79: 6777, Boshart *et al.*, *Cell* (1985) 41: 521 and U.S. 4,399,216. Other features of mammalian expression of metastatic marker expression constructs can be facilitated as described in Ham and Wallace, *Meth. Enz.* (1979) 58: 44, Barnes and Sato, *Anal. Biochem.* (1980) 102: 255, U.S. 4,767,704, US 4,657,866, US 4,927,762, US 4,560,655, WO 90/103430, WO 87/00195, and U.S. RE 30,985.

Subgenomic polynucleotides of the invention can also be used in gene delivery vehicles, for the purpose of delivering a metastatic marker mRNA or of oligonucleotide (either with the sequence of native metastatic marker mRNA or its complement), full-length metastatic marker protein, metastatic marker fusion protein, metastatic marker polypeptide, or metastatic marker-specific ribozyme or single-chain antibody, into a cell preferably a eukaryotic cell. According to the present invention, a gene delivery vehicle can be, for example, naked plasmid DNA, a viral expression vector comprising a metastatic marker subgenomic polynucleotide, or a metastatic marker subgenomic polynucleotide in conjunction with a liposome or a condensing agent.

In one embodiment of the invention, the gene delivery vehicle comprises a promoter and a metastatic marker subgenomic polynucleotide. Preferred promoters are tissue-specific promoters and promoters which are activated by cellular proliferation, such as the thymidine kinase and thymidylate synthase promoters. Other preferred promoters include promoters which are activatable by infection with a virus, such as the α - and β -interferon promoters, and promoters which are activatable by a hormone, such as estrogen. Other promoters which can be used include the Moloney virus LTR, the CMV promoter, and the mouse albumin promoter.

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A metastatic marker gene delivery vehicle can comprise viral sequences such as a viral origin of replication or packaging signal. These viral sequences can be selected from viruses such as astrovirus, coronavirus, orthomyxovirus, papovavirus, paramyxovirus, parvovirus, picornavirus, poxvirus, retrovirus, togavirus or adenovirus.

In a preferred embodiment, the metastatic marker gene delivery vehicle is a recombinant retroviral vector. Recombinant retroviruses and various uses thereof have been described in numerous references including. for example, Mann *et al.*, *Cell 33:*153, 1983, Cane and Mulligan, *Proc. Nat'l Acad. Sci. USA 81:*6349, 1984, Miller *et al.*, *Human Gene Therapy* 1:5-14, 1990, U.S. Patent Nos. 4,405,712, 4,861,719, and 4,980,289. and PCT Application Nos. WO 89/02,468, WO 89/05,349, and WO 90/02,806. Numerous retroviral gene delivery vehicles can be utilized in the present invention, including for example those described in EP 0,415,731; WO 90/07936; WO 94/03622; WO 93/25698; WO 93/25234; U.S. Patent No. 5,219,740; WO 9311230; WO 9310218; Vile and Hart, *Cancer Res.* 53:3860-3864, 1993; Vile and Hart, *Cancer Res.* 53:962-967, 1993; Ram *et al.*, *Cancer Res.* 53:83-88, 1993; Takamiya *et al.*, *J. Neurosci. Res.* 33:493-503, 1992; Baba *et al.*, *J. Neurosurg.* 79:729-735, 1993 (U.S. Patent No. 4,777,127, GB 2,200,651, EP 0,345,242 and WO91/02805).

Particularly preferred retroviruses are derived from retroviruses which include avian leukosis virus (ATCC Nos. VR-535 and VR-247), bovine leukemia virus (VR-1315), murine leukemia virus (MLV), mink-cell focus-inducing virus (Koch et al., J. Vir. 49:828, 1984; and Oliff et al., J. Vir. 48:542, 1983), murine sarcoma virus (ATCC Nos. VR-844, 45010 and 45016), reticuloendotheliosis virus (ATCC Nos VR-994, VR-770 and 45011), Rous sarcoma virus, Mason-Pfizer monkey virus, baboon endogenous virus, endogenous feline retrovirus (e.g., RD114), and mouse or rat gL30 sequences used as a retroviral vector. Particularly preferred strains of MLV from which recombinant retroviruses can be generated include 4070A and 1504A (Hartley and Rowe, J. Vir. 19:19, 1976), Abelson (ATCC No. VR-999), Friend (ATCC No. VR-245), Graffi (Ru et al., J. Vir. 67:4722, 1993; and Yantchev Neoplasma 26:397, 1979). Gross (ATCC No. VR-590), Kirsten (Albino et al., J. Exp. Med. 164:1710, 1986), Harvey sarcoma virus (Manly et al., J. Vir. 62:3540, 1988; and Albino et al., J. Exp. Med. 164:1710, 1986) and Rauscher (ATCC No. VR-998), and Moloney MLV (ATCC No. VR-190). A particularly preferred non-mouse retrovirus is Rous sarcoma virus. Preferred Rous sarcoma viruses include Bratislava (Manly et al., J. Vir. 62:3540, 1988; and Albino et al., J. Exp. Med. 164:1710, 1986), Bryan high titer (e.g., ATCC Nos. VR-

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334, VR-657, VR-726, VR-659, and VR-728), Bryan standard (ATCC No. VR-140), Carr-Zilber (Adgighitov et al., Neoplasma 27:159, 1980), Engelbreth-Holm (Laurent et al., Biochem Biophys Acta 908:241, 1987), Harris, Prague (e.g., ATCC Nos. VR-772, and 45033), and Schmidt-Ruppin (e.g., ATCC Nos. VR-724, VR-725, VR-354) viruses.

Any of the above retroviruses can be readily utilized in order to assemble or construct retroviral metastatic marker gene delivery vehicles given the disclosure provided herein and standard recombinant techniques (e.g., Sambrook et al., Molecular Cloning: A Laboratory Manual, 2d ed., Cold Spring Harbor Laboratory Press, 1989, and Kunkle, PNAS 82:488, 1985) known in the art. Portions of retroviral Metastatic marker expression vectors can be derived from different retroviruses. For example, retrovector LTRs can be derived from a murine sarcoma virus, a tRNA binding site from a Rous sarcoma virus, a packaging signal from a murine leukemia virus, and an origin of second strand synthesis from an avian leukosis virus. These recombinant retroviral vectors can be used to generate transduction competent retroviral vector particles by introducing them into appropriate packaging cell lines (see Serial No. 07/800,921, filed November 29, 1991). Recombinant retroviruses can be produced which direct the site-specific integration of the recombinant retroviral genome into specific regions of the host cell DNA. Such site-specific integration can be mediated by a chimeric integrase incorporated into the retroviral particle (see Serial No. 08/445,466 filed May 22, 1995). It is preferable that the recombinant viral gene delivery vehicle is a replication-defective recombinant virus.

Packaging cell lines suitable for use with the above-described retroviral gene delivery vehicles can be readily prepared (see Serial No. 08/240,030, filed May 9, 1994; see also WO 92/05266) and used to create producer cell lines (also termed vector cell lines or "VCLs") for production of recombinant viral particles. In particularly preferred embodiments of the present invention, packaging cell lines are made from human (e.g., HT1080 cells) or mink parent cell lines, thereby allowing production of recombinant retroviral gene delivery vehicles which are capable of surviving inactivation in human serum. The construction of recombinant retroviral gene delivery vehicles is described in detail in WO 91/02805. These recombinant retroviral gene

delivery vehicles can be used to generate transduction competent retroviral particles by introducing them into appropriate packaging cell lines (*see* Serial No. 07/800,921). Similarly, adenovirus gene delivery vehicles can also be readily prepared and utilized given the disclosure provided herein (*see also* Berkner. *Biotechniques* 6:616-627, 1988, and Rosenfeld *et al.*, *Science* 252:431-434, 1991, WO 93/07283, WO 93/06223, and WO 93/07282).

A metastatic marker gene delivery vehicle can also be a recombinant adenoviral gene delivery vehicle. Such vehicles can be readily prepared and utilized given the disclosure provided herein (see Berkner, Biotechniques 6:616, 1988, and Rosenfeld et al., Science 252:431, 1991, WO 93/07283, WO 93/06223, and WO 93/07282). Adeno-associated viral metastatic marker gene delivery vehicles can also be constructed and used to deliver metastatic marker amino acids or nucleotides. The use of adeno-associated viral gene delivery vehicles in vitro is described in Chatterjee et al., Science 258: 1485-1488 (1992), Walsh et al., Proc. Nat'l Acad. Sci. 89: 7257-7261 (1992), Walsh et al., J. Clin. Invest. 94: 1440-1448 (1994). Flotte et al., J. Biol. Chem. 268: 3781-3790 (1993), Ponnazhagan et al., J. Exp. Med. 179: 733-738 (1994), Miller et al., Proc. Nat'l Acad. Sci. 91: 10183-10187 (1994), Einerhand et al., Gene Ther. 2: 336-343 (1995), Luo et al., Exp. Hematol. 23: 1261-1267 (1995), and Zhou et al., Gene Therapy 3: 223-229 (1996). In vivo use of these vehicles is described in Flotte et al., Proc. Nat'l Acad. Sci. 90: 10613-10617 (1993), and Kaplitt et al., Nature Genet. 8:148-153 (1994).

In another embodiment of the invention, a metastatic marker gene delivery vehicle is derived from a togavirus. Preferred togaviruses include alphaviruses, in particular those described in U.S. Serial No. 08/405,627, filed March 15, 1995, WO 95/07994. Alpha viruses, including Sindbis and ELVS viruses can be gene delivery vehicles for metastatic marker polynucleotides. Alpha viruses are described in WO 94/21792, WO 92/10578 and WO 95/07994. Several different alphavirus gene delivery vehicle systems can be constructed and used to deliver metastatic marker subgenomic polynucleotides to a cell according to the present invention. Representative examples of such systems include those described in U.S. Patents 5,091,309 and 5,217,879.

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Particularly preferred alphavirus gene delivery vehicles for use in the present invention include those which are described in WO 95/07994, and U.S. Serial No. 08/405,627.

Preferably, the recombinant viral vehicle is a recombinant alphavirus viral vehicle based on a Sindbis virus. Sindbis constructs, as well as numerous similar constructs, can be readily prepared essentially as described in U.S. Serial No. 08/198,450. Sindbis viral gene delivery vehicles typically comprise a 5' sequence capable of initiating Sindbis virus transcription, a nucleotide sequence encoding Sindbis non-structural proteins, a viral junction region inactivated so as to prevent subgenomic fragment transcription, and a Sindbis RNA polymerase recognition sequence. Optionally, the viral junction region can be modified so that subgenomic polynucleotide transcription is reduced, increased, or maintained. As will be appreciated by those in the art, corresponding regions from other alphaviruses can be used in place of those described above.

The viral junction region of an alphavirus-derived gene delivery vehicle can comprise a first viral junction region which has been inactivated in order to prevent transcription of the subgenomic polynucleotide and a second viral junction region which has been modified such that subgenomic polynucleotide transcription is reduced. An alphavirus-derived vehicle can also include a '5' promoter capable of initiating synthesis of viral RNA from cDNA and a 3' sequence which controls transcription termination.

Other recombinant togaviral gene delivery vehicles which can be utilized in the present invention include those derived from Semliki Forest virus (ATCC VR-67; ATCC VR-1247), Middleberg virus (ATCC VR-370), Ross River virus (ATCC VR-373; ATCC VR-1246), Venezuelan equine encephalitis virus (ATCC VR923; ATCC VR-1250; ATCC VR-1249; ATCC VR-532), and those described in U.S. Patents 5,091,309 and 5,217,879 and in WO 92/10578. The Sindbis vehicles described above. as well as numerous similar constructs, can be readily prepared essentially as described in U.S. Serial No. 08/198,450.

Other viral gene delivery vehicles suitable for use in the present invention include, for example, those derived from poliovirus (Evans et al., Nature

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339:385, 1989, and Sabin et al., J. Biol. Standardization 1:115, 1973) (ATCC VR-58); rhinovirus (Arnold et al., J. Cell. Biochem. L401, 1990) (ATCC VR-1110); pox viruses, such as canary pox virus or vaccinia virus (Fisher-Hoch et al., PNAS 86:317, 1989; Flexner et al., Ann. N.Y. Acad. Sci. 569:86, 1989; Flexner et al., Vaccine 8:17, 1990; U.S. 4.603.112 and U.S. 4,769,330; WO 89/01973) (ATCC VR-111; ATCC VR-2010); SV40 (Mulligan et al., Nature 277:108, 1979) (ATCC VR-305), (Madzak et al., J. Gen. Vir. 73:1533, 1992); influenza virus (Luytjes et al., Cell 59:1107, 1989; McMicheal et al., The New England Journal of Medicine 309:13, 1983; and Yap et al., Nature 273:238, 1978) (ATCC VR-797); parvovirus such as adeno-associated virus (Samulski et al., J. Vir. 63:3822, 1989, and Mendelson et al., Virology 166:154, 1988) (ATCC VR-645); herpes simplex virus (Kit et al., Adv. Exp. Med. Biol. 215:219, 1989) (ATCC VR-977; ATCC VR-260); Nature 277: 108, 1979); human immunodeficiency virus (EPO 386,882, Buchschacher et al., J. Vir. 66:2731, 1992); measles virus (EPO 440,219) (ATCC VR-24); A (ATCC VR-67; ATCC VR-1247), Aura (ATCC VR-368), Bebaru virus (ATCC VR-600; ATCC VR-1240), Cabassou (ATCC VR-922), Chikungunya virus (ATCC VR-64; ATCC VR-1241), Fort Morgan (ATCC VR-924), Getah virus (ATCC VR-369; ATCC VR-1243), Kyzylagach (ATCC VR-927), Mayaro (ATCC VR-66), Mucambo virus (ATCC VR-580; ATCC VR-1244), Ndumu (ATCC VR-371), Pixuna virus (ATCC VR-372; ATCC VR-1245), Tonate (ATCC VR-925). Triniti (ATCC VR-469). Una (ATCC VR-374), Whataroa (ATCC VR-926), Y-62-33 (ATCC VR-375), O'Nyong virus, Eastern encephalitis virus (ATCC VR-65; ATCC VR-1242), Western encephalitis virus (ATCC VR-70; ATCC VR-1251; ATCC VR-622; ATCC VR-1252), and coronavirus (Hamre et al., Proc. Soc. Exp. Biol. Med. 121:190, 1966) (ATCC VR-740).

A subgenomic metastatic marker polynucleotide of the invention can also be combined with a condensing agent to form a gene delivery vehicle. In a preferred embodiment, the condensing agent is a polycation, such as polylysine, polyarginine, polyornithine, protamine, spermine, spermidine, and putrescine. Many suitable methods for making such linkages are known in the art (see, for example, Serial No. 08/366,787, filed December 30, 1994).

In an alternative embodiment, a metastatic marker subgenomic polynucleotide is associated with a liposome to form a gene delivery vehicle. Liposomes are small, lipid vesicles comprised of an aqueous compartment enclosed by a lipid bilayer, typically spherical or slightly elongated structures several hundred Angstroms in diameter. Under appropriate conditions, a liposome can fuse with the plasma membrane of a cell or with the membrane of an endocytic vesicle within a cell which has internalized the liposome, thereby releasing its contents into the cytoplasm. Prior to interaction with the surface of a cell, however, the liposome membrane acts as a relatively impermeable barrier which sequesters and protects its contents, for example, from degradative enzymes. Additionally, because a liposome is a synthetic structure, specially designed liposomes can be produced which incorporate desirable features. See Stryer, Biochemistry, pp. 236-240, 1975 (W.H. Freeman, San Francisco, CA); Szoka et al., Biochim. Biophys. Acta 600:1, 1980; Bayer et al., Biochim. Biophys. Acta. 550:464, 1979; Rivnay et al., Meth. Enzymol. 149:119, 1987; Wang et al., PNAS 84: 7851, 1987, Plant et al., Anal. Biochem. 176:420, 1989, and U.S. Patent 4,762,915. Liposomes can encapsulate a variety of nucleic acid molecules including DNA, RNA, plasmids, and expression constructs comprising metastatic marker subgenomic polynucleotides such those disclosed in the present invention.

Liposomal preparations for use in the present invention include cationic (positively charged), anionic (negatively charged) and neutral preparations. Cationic liposomes have been shown to mediate intracellular delivery of plasmid DNA (Felgner et al., Proc. Nat'l Acad. Sci. USA 84:7413-7416, 1987), mRNA (Malone et al., Proc. Nat'l Acad. Sci. USA 86:6077-6081, 1989), and purified transcription factors (Debs et al., J. Biol. Chem. 265:10189-10192, 1990), in functional form. Cationic liposomes are readily available. For example, N[1-2,3-dioleyloxy)propyl]-N,N.N-triethylammonium (DOTMA) liposomes are available under the trademark Lipofectin, from GIBCO BRL, Grand Island, NY. See also Felgner et al., Proc. Nat'l Acad. Sci. USA 91: 5148-5152.87, 1994. Other commercially available liposomes include Transfectace (DDAB/DOPE) and DOTAP/DOPE (Boerhinger). Other cationic liposomes can be prepared from readily available materials using techniques well known in the art. See,

e.g., Szoka et al., Proc. Nat'l Acad. Sci. USA 75:4194-4198, 1978; and WO 90/11092 for descriptions of the synthesis of DOTAP (1,2-bis(oleoyloxy)-3-(trimethylammonio)propane) liposomes.

Similarly, anionic and neutral liposomes are readily available, such as from Avanti Polar Lipids (Birmingham, AL), or can be easily prepared using readily available materials. Such materials include phosphatidyl choline, cholesterol, phosphatidyl ethanolamine, dioleoylphosphatidyl choline (DOPC), dioleoylphosphatidyl glycerol (DOPG), dioleoylphoshatidyl ethanolamine (DOPE), among others. These materials can also be mixed with the DOTMA and DOTAP starting materials in appropriate ratios. Methods for making liposomes using these materials are well known in the art.

The liposomes can comprise multilammelar vesicles (MLVs), small unilamellar vesicles (SUVs), or large unilamellar vesicles (LUVs). The various liposome-nucleic acid complexes are prepared using methods known in the art. See, e.g., Straubinger et al., METHODS OF IMMUNOLOGY (1983), Vol. 101, pp. 512-527; Szoka et al., Proc. Nat'l Acad. Sci. USA 87:3410-3414, 1990; Papahadjopoulos et al., Biochim. Biophys. Acta 394:483, 1975; Wilson et al., Cell 17:77, 1979; Deamer and Bangham, Biochim. Biophys. Acta 443:629, 1976; Ostro et al., Biochem. Biophys. Res. Commun. 76:836, 1977; Fraley et al., Proc. Nat'l Acad. Sci. USA 76:3348, 1979; Enoch and Strittmatter, Proc. Nat'l Acad. Sci. USA 76:145, 1979; Fraley et al., J. Biol. Chem. 255:10431, 1980; Szoka and Papahadjopoulos, Proc. Nat'l Acad. Sci. USA 75:145, 1979; and Schaefer-Ridder et al., Science 215:166, 1982.

In addition, lipoproteins can be included with a metastatic marker subgenomic polynucleotide for delivery to a cell. Examples of such lipoproteins include chylomicrons, HDL, IDL, LDL, and VLDL. Mutants. fragments, or fusions of these proteins can also be used. Modifications of naturally occurring lipoproteins can also be used, such as acetylated LDL. These lipoproteins can target the delivery of polynucleotides to cells expressing lipoprotein receptors. Preferably, if lipoproteins are included with a polynucleotide, no other targeting ligand is included in the composition.

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In another embodiment, naked metastatic marker subgenomic polynucleotide molecules are used as gene delivery vehicles, as described in WO 90/11092 and U.S. Patent 5,580,859. Such gene delivery vehicles can be either metastatic marker DNA or RNA and, in certain embodiments, are linked to killed adenovirus. Curiel et al., Hum. Gene. Ther. 3:147-154, 1992. Other suitable vehicles include DNA-ligand (Wu et al., J. Biol. Chem. 264:16985-16987, 1989), lipid-DNA combinations (Felgner et al., Proc. Nat'l Acad. Sci. USA 84:7413 7417, 1989), liposomes (Wang et al., Proc. Nat'l Acad. Sci. 84:7851-7855, 1987) and microprojectiles (Williams et al., Proc. Nat'l Acad. Sci. 88:2726-2730, 1991).

One can increase the efficiency of naked metastatic marker subgenomic polynucleotide uptake into cells by coating the polynucleotides onto biodegradable latex beads. This approach takes advantage of the observation that latex beads, when incubated with cells in culture, are efficiently transported and concentrated in the perinuclear region of the cells. The beads will then be transported into cells when injected into muscle. Metastatic marker subgenomic polynucleotide-coated latex beads will be efficiently transported into cells after endocytosis is initiated by the latex beads and thus increase gene transfer and expression efficiency. This method can be improved further by treating the beads to increase their hydrophobicity, thereby facilitating the disruption of the endosome and release of metastatic marker subgenomic polynucleotides into the cytoplasm.

The invention provides a method of detecting metastatic marker gene expression in a biological sample. Detection of metastatic marker gene expression is useful, for example, for identifying metastases or for determining metastatic potential in a tissue sample, preferably a tumor. Appropriate treatment regimens can then be designed for patients who are at risk for developing metastatic cancers in other organs of the body.

The body sample can be, for example, a solid tissue or a fluid sample. Protein or nucleic acid expression products can be detected in the body sample. In one embodiment, the body sample is assayed for the presence of a metastatic marker protein. A metastatic marker protein comprises a sequence encoded by a nucleotide

sequence shown in SEQ ID NOS:1-85 or its complement and can be detected using the marker protein-specific antibodies of the present invention. The antibodies can be labeled, for example, with a radioactive, fluorescent, biotinylated, or enzymatic tag and detected directly, or can be detected using indirect immunochemical methods, using a labeled secondary antibody. The presence of the metastatic marker proteins can be assayed, for example, in tissue sections by immunocytochemistry, or in lysates, using Western blotting, as is known in the art.

In another embodiment, the body sample is assayed for the presence of marker protein mRNA. A sample can be contacted with a nucleic acid hybridization probe capable of hybridizing with the mRNA corresponding the selected polypeptide. Still further, the sample can be subjected to a Northern blotting technique to detect mRNA, indicating expression of the polypeptide. For those techniques in which mRNA is detected, the sample can be subjected to a nucleic acid amplification process whereby the mRNA molecule or a selected part thereof is amplified using appropriate nucleotide primers. Other RNA detection techniques can also be used, including, but not limited to, *in situ* hybridization.

Marker protein-specific probes can be generated using the cDNA sequences disclosed in SEQ ID NOS:1-85. The probes are preferably at least 15 to 50 nucleotides in length, although they can be at least 8, 10, 11, 12, 20, 25, 30, 35, 40, 45, 60, 75, or 100 or more nucleotides in length. The probes can be synthesized chemically or can be generated from longer polynucleotides using restriction enzymes. The probes can be labeled, for example, with a radioactive, biotinylated, or fluorescent tag.

Optionally, the level of a particular metastatic marker expression product in a body sample can be quantitated. Quantitation can be accomplished, for example, by comparing the level of expression product detected in the body sample with the amounts of product present in a standard curve. A comparison can be made visually or using a technique such as densitometry, with or without computerized assistance. For use as controls, body samples can be isolated from other humans, other non-cancerous organs of the patient being tested, or non-metastatic breast or colon cancer from the patient being tested.

Polynucleotides encoding metastatic marker-specific reagents of the invention, such as antibodies and nucleotide probes, can be supplied in a kit for detecting marker gene expression products in a biological sample. The kit can also contain buffers or labeling components, as well as instructions for using the reagents to detect the marker expression products in the biological sample.

If expression of a metastatic marker gene having a nucleotide sequence shown in SEQ ID NOS:2, 4, 9, 13, 14, 19, 26, 29, 39-41, 48, 55, 57, 60, 63, 64, 72, 73, 82, or 83 is detected, the biological sample contains cancer cells which will likely metastasize to the lung. If expression of a metastatic marker gene having a nucleotide sequence shown in SEQ ID NOS:1, 5, 11, 18, 20, 22, 24, 30, 33, 35, 36, 38, 45, 52, 58. 65, 66, 70, 74, 76, or 80 is detected, the biological sample contains cancer cells which will likely metastasize to the bone and/or lung. On the other hand, if expression of a metastatic marker gene having a nucleotide sequence shown in SEQ ID NOS:3, 7, 8, 10, 12, 15-17, 21, 23, 25, 28, 31, 34, 37, 42-44, 46, 47, 49-51, 53, 59, 61, 62, 67, 68, 75, 77-79, 81, 84, or 85 is detected, the biological sample contains cancer cells which will likely not metastasize. Detection of expression of a metastatic marker gene comprising the nucleotide sequence shown in SEQ ID NO:56 also indicates that the biological sample contains cancer cells which will likely metastasize. This information can be used, for example, to design treatment regimens. Treatment regiments can include altering expression of one or more metastatic marker genes, as desired. Metastatic marker gene expression can be altered for therapeutic purposes, as described below, or can be used to identify therapeutic agents.

In one embodiment of the invention, expression of a metastatic marker gene whose expression is up-regulated in metastatic cancer is decreased using a ribozyme, an RNA molecule with catalytic activity. See, e.g., Cech, 1987, Science 236: 1532-1539; Cech, 1990, Ann. Rev. Biochem. 59:543-568; Cech. 1992, Curr. Opin. Struct. Biol. 2: 605-609; Couture and Stinchcomb. 1996, Trends Genet. 12: 510-515. Ribozymes can be used to inhibit gene function by cleaving an RNA sequence. as is known in the art (e.g., Haseloff et al., U.S. 5,641.673).

Coding sequences of metastatic marker genes can be used to generate ribozymes which will specifically bind to mRNA transcribed from a metastatic marker gene. Methods of designing and constructing ribozymes which can cleave other RNA molecules in trans in a highly sequence specific manner have been developed and described in the art (see Haseloff, J. et al. (1988), Nature 334:585-591). For example, the cleavage activity of ribozymes can be targeted to specific RNAs by engineering a discrete "hybridization" region into the ribozyme. The hybridization region contains a sequence complementary to the target RNA and thus specifically hybridizes with the target (see, for example, Gerlach, W. L. et al., EP 321,201). Longer complementary sequences can be used to increase the affinity of the hybridization sequence for the target. The hybridizing and cleavage regions of the ribozyme can be integrally related; thus, upon hybridizing to the target RNA through the complementary regions, the catalytic region of the ribozyme can cleave the target.

Ribozymes can be introduced into cells as part of a DNA construct, as is known in the art. The DNA construct can also include transcriptional regulatory elements, such as a promoter element, an enhancer or UAS element, and a transcriptional terminator signal, for controlling the transcription of the ribozyme in the cells.

Mechanical methods, such as microinjection. liposome-mediated transfection, electroporation, or calcium phosphate precipitation, can be used to introduce a ribozyme-containing DNA construct into cells whose division it is desired to decrease, as described above. Alternatively, if it is desired that a DNA construct be stably retained by the cells, the DNA construct can be supplied on a plasmid and maintained as a separate element or integrated into the genome of the cells, as is known in the art.

As taught in Haseloff *et al.*, U.S. 5,641.673. ribozymes can be engineered so that their expression will occur in response to factors which induce expression of metastatic marker genes. Ribozymes can also be engineered to provide an additional level of regulation, so that destruction of mRNA occurs only when both a ribozyme and a metastatic marker gene are expressed in the cells.

Expression of a metastatic marker gene can also be altered using an antisense oligonucleotide sequence. The antisense sequence is complementary to at least a portion of the coding sequence of a metastatic marker gene having a nucleotide sequence shown in SEQ ID NOS: 1-85. The complement of a nucleotide sequence shown in SEQ ID NOS: 1-85 is a contiguous sequence of nucleotides which form Watson-Crick basepairs with a contiguous nucleotide sequence shown in SEQ ID NOS: 1-85.

Preferably, the antisense oligonucleotide sequence is at least six nucleotides in length, but can be at least about 8, 12, 15, 20, 25, 30, 35, 40, 45, or 50 nucleotides long. Longer sequences can also be used. Antisense oligonucleotide molecules can be provided in a DNA construct and introduced into cells whose division is to be decreased, as described above.

Antisense oligonucleotides can comprise deoxyribonucleotides, ribonucleotides, or a combination of both. Oligonucleotides can be synthesized manually or by an automated synthesizer, by covalently linking the 5' end of one nucleotide with the 3' end of another nucleotide with non-phosphodiester internucleotide linkages such alkylphosphonates, phosphorothioates, phosphorodithioates, alkylphosphonothioates, alkylphosphonates, phosphoramidates, phosphate esters, carbamates, acetamidate, carboxymethyl esters, carbonates, and phosphate triesters. See Brown, 1994, Meth. Mol. Biol. 20:1-8; Sonveaux, 1994, Meth. Mol. Biol. 26:1-72; Uhlmann et al., 1990, Chem. Rev. 90:543-583.

Although precise complementarity is not required for successful duplex formation between an antisense molecule and the complementary coding sequence of a metastatic marker gene, antisense molecules with no more than one mismatch are preferred. One skilled in the art can easily use the calculated melting point of a metastatic marker gene antisense-sense pair to determine the degree of mismatching which will be tolerated between a particular antisense oligonucleotide and a particular coding sequence of the selected gene.

Antisense oligonucleotides can be modified without affecting their ability to hybridize to a metastatic marker protein coding sequence. These

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modifications can be internal or at one or both ends of the antisense molecule. For example, internucleoside phosphate linkages can be modified by adding cholesteryl or diamine moieties with varying numbers of carbon residues between the amino groups and terminal ribose. Modified bases and/or sugars, such as arabinose instead of ribose, or a 3', 5'-substituted oligonucleotide in which the 3' hydroxyl group or the 5' phosphate group are substituted, can also be employed in a modified antisense oligonucleotide. These modified oligonucleotides can be prepared by methods well known in the art. Agrawal et al., 1992, Trends Biotechnol. 10:152-158; Uhlmann et al., 1990, Chem. Rev. 90:543-584; Uhlmann et al., 1987, Tetrahedron. Lett. 215:3539-3542.

Antibodies of the invention which specifically bind to a metastatic marker protein can also be used to alter metastatic marker gene expression. By antibodies is meant antibodies and parts or derivatives thereof, such as single chain antibodies, that retain specific binding for the protein. Specific antibodies bind to metastatic marker proteins and prevent the proteins from functioning in the cell. Polynucleotides encoding specific antibodies of the invention can be introduced into cells, as described above.

Marker proteins of the present invention can be used to screen for drugs which have a therapeutic anti-metastatic effect. The effect of a test compound on metastatic marker protein synthesis can also be used to identify test compounds which modulate metastasis. Test compounds which can be screened include any substances, whether natural products or synthetic, which can be administered to the subject. Libraries or mixtures of compounds can be tested. The compounds or substances can be those for which a pharmaceutical effect is previously known or unknown.

A cell is contacted with a test compound. The cell can be any cell, such as a colon cancer cell, which ordinarily synthesizes the metastatic marker protein being measured. For example, Tables 1 and 2 provide appropriate cell types which can be used for screening assays.

Synthesis of metastatic marker proteins can be measured by any means for measuring protein synthesis known in the art, such as incorporation of labeled amino acids into proteins and detection of labeled metastatic marker proteins in a

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polyacrylamide gel. The amount of metastatic marker proteins can be detected, for example, using metastatic marker protein-specific antibodies of the invention in Western blots. The amount of the metastatic marker proteins synthesized in the presence or absence of a test compound can be determined by any means known in the art, such as comparison of the amount of metastatic marker protein synthesized with the amount of the metastatic marker proteins present in a standard curve.

The effect of a test compound on metastatic marker protein synthesis can also be measured by Northern blot analysis, by measuring the amount of metastatic marker protein mRNA expression in response to the test compound using metastatic marker protein specific nucleotide probes of the invention, as is known in the art.

Typically, biological sample is contacted with a range of concentrations of the test compound, such as 1.0 nM, 5.0 nM, 10 nM, 50 nM, 100 nM, 500 nM, 1 mM, 10 mM, 50 mM, and 100 mM. Preferably, the test compound increases or decreases expression of a metastatic marker protein by 60%. 75%, or 80%. More preferably, an increase or decrease of 85%, 90%, 95%, or 98% is achieved.

The invention provides compositions for increasing or decreasing expression of metastatic marker protein. Therapeutic compositions for increasing metastatic marker gene expression are desirable for markers which are down-regulated in metastatic cells. These compositions comprise polynucleotides encoding all or at least a portion of a metastatic marker protein gene expression product. Preferably, the therapeutic composition contains an expression construct comprising a promoter and a polynucleotide segment encoding at least a portion of the metastatic marker protein which is effective to increase or decrease metastatic potential. Portions of metastatic marker genes or proteins which are effective to decrease metastatic potential of a cell can be determined, for example, by introducing various portions of metastatic marker genes or polypeptides into metastatic cell lines, such as MDA-MB-231, MDA-MB-435, Km12C, or Km12L4, and assaying the division rate of the cells or the ability of the cells to form metastases when implanted *in vivo*, as is known in the art. Non-metastatic cell lines, such as MCF-7, can be used to assay the ability of a portion of a metastatic marker protein to increase expression of a metastatic marker gene.

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Within the expression construct, the polynucleotide segment is located downstream from the promoter, and transcription of the polynucleotide segment initiates at the promoter. A more complete description of gene transfer vectors, especially retroviral vectors is contained in U.S. Serial No. 08/869,309, which is incorporated herein by reference.

Decreased metastatic marker gene expression is desired in conditions in which the marker gene is up-regulated in metastatic cancer. Therapeutic compositions for treating these disorders comprise a polynucleotide encoding a reagent which specifically binds to a metastatic marker protein expression product, as disclosed herein.

Metastatic marker therapeutic compositions of the invention can comprise a pharmaceutically acceptable carrier. Pharmaceutically acceptable carriers are well known to those in the art. Such carriers include, but are not limited to, large, slowly metabolized macromolecules, such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and inactive virus particles. Pharmaceutically acceptable salts can also be used in the composition, for example, mineral salts such as hydrochlorides, hydrobromides, phosphates, or sulfates, as well as the salts of organic acids such as acetates, proprionates, malonates, or benzoates.

Therapeutic compositions can also contain liquids, such as water, saline, glycerol, and ethanol, as well as substances such as wetting agents, emulsifying agents, or pH buffering agents. Liposomes, such as those described in U.S. 5,422,120, WO 95/13796, WO 91/14445, or EP 524,968 B1, can also be used as a carrier for the therapeutic composition.

Typically, a therapeutic metastatic marker composition is prepared as an injectable, either as a liquid solution or suspension; however, solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection can also be prepared. A metastatic marker composition can also be formulated into an enteric coated tablet or gel capsule according to known methods in the art, such as those described in U.S. 4,853,230, EP 225,189, AU 9,224,296, and AU 9,230,801.

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Administration of the metastatic marker therapeutic agents of the invention can include local or systemic administration, including injection, oral administration, particle gun, or catheterized administration, and topical administration. Various methods can be used to administer a therapeutic metastatic marker composition directly to a specific site in the body.

For treatment of tumors, including metastatic lesions, for example, a therapeutic metastatic marker composition can be injected several times in several different locations within the body of tumor. Alternatively, arteries which serve a tumor can be identified, and a therapeutic composition injected into such an artery, in order to deliver the composition directly into the tumor.

A tumor which has a necrotic center can be aspirated and the composition injected directly into the now empty center of the tumor. A therapeutic metastatic marker composition can be directly administered to the surface of a tumor, for example, by topical application of the composition. X-ray imaging can be used to assist in certain of the above delivery methods. Combination therapeutic agents, including a metastatic marker proteins or polypeptide or a metastatic marker subgenomic polynucleotide and other therapeutic agents, can be administered simultaneously or sequentially.

Receptor-mediated targeted delivery can be used to deliver therapeutic compositions containing metastatic marker subgenomic polynucleotides, proteins, or reagents such as antibodies, ribozymes, or antisense oligonucleotides to specific tissues. Receptor-mediated delivery techniques are described in, for example, Findeis et al. (1993), *Trends in Biotechnol. 11*, 202-05; Chiou et al. (1994), GENE THERAPEUTICS: METHODS AND APPLICATIONS OF DIRECT GENE TRANSFER (J.A. Wolff, ed.); Wu & Wu (1988), *J. Biol. Chem. 263*, 621-24; Wu et al. (1994). *J. Biol. Chem. 269*, 542-46; Zenke et al. (1990), *Proc. Nat'l Acad. Sci. U.S.A. 87*, 3655-59; Wu et al. (1991), *J. Biol. Chem. 266*, 338-42.

Alternatively, a metastatic marker therapeutic composition can be introduced into human cells *ex vivo*, and the cells then replaced into the human. Cells can be removed from a variety of locations including, for example, from a selected

tumor or from an affected organ. In addition, a therapeutic composition can be inserted into non-affected, for example, dermal fibroblasts or peripheral blood leukocytes. If desired, particular fractions of cells such as a T cell subset or stem cells can also be specifically removed from the blood (see, for example, PCT WO 91/16116). The removed cells can then be contacted with a metastatic marker therapeutic composition utilizing any of the above-described techniques, followed by the return of the cells to the human, preferably to or within the vicinity of a tumor or other site to be treated. The methods described above can additionally comprise the steps of depleting fibroblasts or other non-contaminating tumor cells subsequent to removing tumor cells from a human, and/or the step of inactivating the cells, for example, by irradiation.

Both the dose of a metastatic marker composition and the means of administration can be determined based on the specific qualities of the therapeutic composition, the condition, age, and weight of the patient, the progression of the disease, and other relevant factors. Preferably, a therapeutic composition of the invention increases or decreases expression of the metastatic marker genes by 50%, 60%, 70%, or 80%. Most preferably, expression of the metastatic marker genes is increased or decreased by 90%, 95%, 99%, or 100%. The effectiveness of the mechanism chosen to alter expression of the metastatic marker genes can be assessed using methods well known in the art, such as hybridization of nucleotide probes to mRNA of the metastatic marker genes, quantitative RT-PCR, or detection of an the metastatic marker proteins using specific antibodies of the invention.

If the composition contains the metastatic marker proteins, polypeptide, or antibody, effective dosages of the composition are in the range of about 5 μ g to about 50 μ g/kg of patient body weight, about 50 μ g to about 5 mg/kg, about 100 μ g to about 500 μ g/kg of patient body weight, and about 200 to about 250 μ g/kg.

Therapeutic compositions containing metastatic marker subgenomic polynucleotides can be administered in a range of about 100 ng to about 200 mg of DNA for local administration. Concentration ranges of about 500 ng to about 50 mg, about 1 µg to about 2 mg, about 5 µg to about 500 µg, and about 20 µg to about 100 µg of DNA can also be used during a gene therapy protocol. Factors such as method of

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action and efficacy of transformation and expression are considerations that will affect the dosage required for ultimate efficacy of the metastatic marker subgenomic polynucleotides. Where greater expression is desired over a larger area of tissue, larger amounts of metastatic marker subgenomic polynucleotides or the same amounts readministered in a successive protocol of administrations, or several administrations to different adjacent or close tissue portions of, for example, a tumor site, can be required to effect a positive therapeutic outcome. In all cases, routine experimentation in clinical trials will determine specific ranges for optimal therapeutic effect.

Expression of an endogenous metastatic marker gene in a cell can also be altered by introducing in frame with the endogenous metastatic marker gene a DNA construct comprising a metastatic marker protein targeting sequence, a regulatory sequence, an exon, and an unpaired splice donor site by homologous recombination, such that a homologously recombinant cell comprising the DNA construct is formed. The new transcription unit can be used to turn the metastatic marker gene on or off as desired. This method of affecting endogenous gene expression is taught in U.S. Patent No. 5,641,670, which is incorporated herein by reference.

The targeting sequence is a segment of at least 10, 12, 15, 20, or 50 contiguous nucleotides selected from the nucleotide sequences shown in SEQ ID NOS:1-85 or the complements thereof. The transcription unit is located upstream of a coding sequence of the endogenous metastatic marker protein gene. The exogenous regulatory sequence directs transcription of the coding sequence of the metastatic marker genes.

A metastatic marker subgenomic polynucleotide can also be delivered to subjects for the purpose of screening test compounds for those which are useful for enhancing transfer of metastatic marker subgenomic polynucleotides to the cell or for enhancing subsequent biological effects of metastatic marker subgenomic polynucleotides within the cell. Such biological effects include hybridization to complementary metastatic marker mRNA and inhibition of its translation, expression of a metastatic marker subgenomic polynucleotide to form metastatic marker mRNA and/or metastatic marker protein, and replication and integration of a metastatic marker

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subgenomic polynucleotide. The subject can be a cell culture or an animal, preferably a mammal, more preferably a human.

Test compounds which can be screened include any substances, whether natural products or synthetic, which can be administered to the subject. Libraries or mixtures of compounds can be tested. The compounds or substances can be those for which a pharmaceutical effect is previously known or unknown. The compounds or substances can be delivered before, after, or concomitantly with a metastatic marker subgenomic polynucleotide. They can be administered separately or in admixture with a metastatic marker subgenomic polynucleotide.

Integration of a delivered metastatic marker subgenomic polynucleotide can be monitored by any means known in the art. For example, Southern blotting of the delivered metastatic marker subgenomic polynucleotide can be performed. A change in the size of the fragments of a delivered polynucleotide indicates integration. Replication of a delivered polynucleotide can be monitored *inter alia* by detecting incorporation of labeled nucleotides combined with hybridization to a metastatic marker probe. Expression of metastatic marker subgenomic polynucleotide can be monitored by detecting production of metastatic marker mRNA which hybridizes to the delivered polynucleotide or by detecting metastatic marker protein. Metastatic marker protein can be detected immunologically. Thus, the delivery of metastatic marker subgenomic polynucleotides according to the present invention provides an excellent system for screening test compounds for their ability to enhance transfer of metastatic marker subgenomic polynucleotides to a cell, by enhancing delivery, integration, hybridization, expression, replication or integration in a cell *in vitro* or in an animal, preferably a mammal, more preferably a human.

The above disclosure generally describes the present invention. A more complete understanding can be obtained by reference to the following specific examples which are provided herein for purposes of illustration only, and are not intended to limit the scope of the invention.

EXAMPLE 1

DIFFERENTIALLY EXPRESSED GENES

This example demonstrates polynucleotides that are differentially expressed in human breast or colon cancer cell lines.

Human cell lines used to identify differentially expressed polynucleotides are the human breast cancer cell lines MCF-7 (non-metastatic), MDA-MB-231 (metastatic to bone and/or lung), and MDA-MB-435 (metastatic to lung) (Brinkley and Cailleau, 1980, *Cancer Res. 40*:3118), and the colon cancer cell lines Km12C (low metastatic) and Km12L4A (highly metastatic) (Morikawa *et al.*, 1988, *Cancer Res. 48*:1943-1948).

RNA was prepared from each cell line and reverse transcribed to form cDNA. The cDNA was amplified using random primers. Amplification products were visualized on a sequencing gel, and cDNA corresponding to mRNA which was differentially expressed in the cell lines was identified.

Expression patterns and sequence identification numbers of novel metastatic marker polynucleotides are shown in Table 1.

Expression patterns and sequence identification numbers of metastatic marker polynucleotides which correspond to known genes are shown in Table 2, and the corresponding proteins are described below.

Osteopontin (SEQ ID NO:64) (OPN or Spp1 for secreted phosphoprotein 1) is a secreted extracellular matrix protein, often expressed during wound healing, involved in osteoclastic differentiation and activation, as described in Heymann *et al.*, 1998, *Cytokine 10*:155-168. Osteopontin is found in bone and other epithelial cells, and has been shown to stimulate proliferation of a quiescent subpopulation of prostate epithelial cells (see Elgavish *et al.*, 1998, *Prostate 35*:83-94).

Osteopontin is implicated during the development of diabetic nephropathy (Fischer et al., 1998, Diabetes 47:1512-1518); the process of cartilage-to-bone transition during rigid bone healing after bone fracture (Nakase et al., 1998, Acta Histochem 100:287-295); wound healing by an interaction with the receptor integrin

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alpha(v)beta 3 after focal stroke (Ellison et al., 1998, Stroke 29:1698-1706); integrin receptor binding and signaling during cell attachment and mechanical stimulation of osteoblasts (Carvalho et al., 1998, J. Cell Biochem 70:376-390); kidney morphogenesis (Denda et al., 1998, Mol. Biol. Cell 9:1425-1435); and as an interstitial chemoattractant in renal inflammation (Rovin and Phan, 1998, Am. J. Kidney Dis. 31:1065-1084). Mice lacking the osteopontin gene showed modulation in osteoclast differentiation from wild type mice (see Rittling et al., 1998, J. Bone Miner Res. 13:1101-1111).

Osteopontin is synthesized by monocytes and macrophages within injury sites, and can promote leukocyte adhesion through the alpha 4beta1 integrin, as described in Bayless *et al.*, 1998, *J. Cell Sci. 111*:1165-1174. Osteopontin is transcriptionally regulated by retinoic acid (see Manji *et al.*, 1998, *J. Cell Physiol. 176*:1-9); preferentially expressed in high grade metastatic brain tumors compared to low grade brain tumors, and inducible by tissue plasminogen activator (tPA) in glioma cell lines (see Tucker *et al.*, 1998, *Anticancer Res. 18*:807-812). Osteopontin is expressed in about 73% of primary gastric carcinoma tissues and correlated with the progression of human gastric carcinoma and lymphogenous metastasis (see Ue *et al.*, 1998, *Int. J. Cancer 79*:127-132).

Nip (SEQ ID NO:65) is described in Boyd et al., 1994, Cell 79:341-351. Adenovirus E1B 19 kDa protein protects against cell death induced by viral infection and external stimuli, and can be functionally substituted with the Bcl-2 protoncogene. E1B 19 kDa interacting proteins (Nip1, Nip2, and Nip3) were discovered in yeast two-hybrid studies conducted to discern proteins that interact with 19 kDa protein, as described by Boyd et al., supra. Nip 1, 2, and 3 interact with discrete domains of E1B 19 kDa, and similarly also interact with Bcl-2, in both cases promoting cell survival.

<u>Ca-dependent protease</u> (SEQ ID NO:66) is Ca⁻²-dependent protease (also called calpain), activity of which is present in every vertebrate cell that has been examined. Ca⁻²-dependent protease activity is associated with cleavages that alter regulation of various enzyme activities, with remodeling or disassembly of the cell cytoskeleton, and with cleavages of hormone receptors (see Goll *et al.*, 1992, *Bioessays* 14(8):549-556). Ca⁻²-dependent protease activity is regulated by binding of Ca⁻² to

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specific sites on the calpain molecule, with binding to each site generating a specific response corelated with a specific activity (e.g., proteolytic activity, calpastatin binding. etc.), as described in Goll *et al*. Excessive activation of the Ca⁺²-dependent protease calpain may play a role in the pathology of disorders including cerebral ischemia, cataract, myocardial ischemia, muscular dystrophy, and platelet aggregation. Therapeutic applications include selective Ca⁻²-dependent protease inhibition, as described in Wang and Yuen, 1994, *Trends Pharmacol. Sci. 15*(11):412-419.

IGF-R (insulin-like growth factor receptor) (SEQ ID NO:67) is a transmembrane tyrosine kinase linked to the ras-raf-MAPK(mitogen-activated protein kinase) cascade and required for the cell to progress through the cell cycle (Werner and Roith. 1997. *Crit. Rev. Oncog* 8(1):71-92). IGF-R mediates mitogenesis, growth hormone action, cell survival and transformation to and maintenance of the malignant phenotype. IGF-R is a member of the growth factor receptor tyrosine kinase superfamily, exists as covalent cross-linked dimers where each monomer is composed of two subunits, and is bound by ligand in the extracellular domain (McInnes and Sykes, 1997, *Biopolymers* 43(5):339-366).

The domains of the IGF-R are described in Sepp-Lorenzino, 1998, Breast Cancer Res Treat 47(3):235-253, including domains responsible for mitogenesis, transformation, and protection from apoptosis. IGF-R expression is increased in breast cancer cells derived from tumor tissue and cell lines, as described in Surmacz et al., 1998, Breast Cancer Res Treat 47(3):255-267, and increased IGF-R may increase tumor mass and/or aid tumor recurrence by promoting proliferation, cell survival, and cell-cell interactions. Human pancreatic cancers overexpress IGF-R and its ligand (Korc, 1998, Surg Oncol Clin N Am 7(1): 25-41), and expression of IGF-I and IGF-R is determined to be a prognostic factor (reflecting the interaction between the neoplastic cells and their microenvironment) for lymphocytic infiltration in thryoid carcinomas (Fonseca et al., 1997, Verh Dtsch Ges Pathol 81:82-96).

ILGF-BP5 (SEQ ID NO:68) is insulin-like growth factor binding protein 5, described in Allander *et al.*, 1994, *J. Biol. Chem. 269*:10891-10898. The gene and promoter for IGF-BP5 are characterized by Allander *et al.*, 1994, *J. Biol Chem.*

269:10891-10898, and some general actions of IGF-BPs are described in Chan and Spencer, 1997, Endocrine 7:95-97. Potential impact of IGF-BPs on cancer cell growth is described in Oh, 1997, Endocrine 7:111-113, and Oh, 1998, Breast Cancer Res Treat 47:283-293. IGF-BP5 is expressed during brain development: IGF-BP5 and IGF-1 mRNAs are synchronously coexpressed in principal neurons of sensory relay systems, including the olfactory bulb, medial and dorsal lateral geniculate bodies, and ventral tier, cochlear, lemniscal, and vestibular nuclei, and are transiently coexpressed in principal neurons of the anterodorsal nucleus, as described in Bondy and Lee, 1993, J. Neurosci 13(12):5092-5104. IGF-BP5 is expressed by luminal or cumulus granulosa cells in virtually all follicles, and is highly abundant in stromal interstitial cells of the mature ovary (see Zhou and Bondy, 1993, Biol. Reprod 48:467-482). IGF-BP5 induction is strongly stimulated during differentiation of skeletal myoblasts and is correlated with IGF-R activation as described in Rousse et al., 1998, Endocrinology 139:1487-1493. IGF-BP5 and other components of the IGF system are critical in postnatal brain development (see Lee et al., 1996, J. Cereb Blood Flow Metab 16:227-236).

IGF-BP5 stimulates bone cell proliferation by an IGF-independent mechanism involving IGF-BP5-specific cell surface binding sites, as described in Mohan *et al.*, 1995, *J. Biol Chem 270*:20424-20431. In connective tissue cell types, IGF-BP5 has a lowered binding affinity to the extracellular matrix which allows IGF-I to better equilibrate with the receptors which in turn potentiates IGF-I action on fibroblasts and smooth muscle cells (Clemmons, *Mol Cell Endocrinology 140*:19-24).

Lactate dehydrogenase (SEQ ID NO:69) is a member of the LDH group of tetrameric enzymes with five isoforms composed of combinations of two subunits. LDH-A and LDH-B. Shim *et al.*, 1997, *Proc. Nat'l Acad. Sci. 94*:6658-6663, described the relationship between LDH-A and neoplasia. In particular, overexpression on LDH-A may contribute to altered metabolism that confers neoplastic growth advantage. The expression pattern of LDH in the present invention is consistent, in that LDH expression is higher in two metastatic breast cancer cell lines than in a non-metastatic breast cancer cell line (Table 2). High or increasing lactate dehydrogenase (LDH) levels

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in tumor tissue and cells is associated with poor survival rate in small cell lung carcinoma (SCLC), as described in Ray et al., 1998. Cancer Detect Prev 22:293-304, making it a useful prognostic indicator for SCLC as discussed in Stokkel et al., 1998, J. Cancer Res Clin Oncol 124:215-219.

<u>Ufo TKR</u> (SEQ ID NO:70) is described in Schulz *et al.*, 1993, *Oncogene* 8:509-513. This protein has been reported as a marker in tumors, but has not previously been reported in breast cancer. According to the present invention, expression is found in the MDA-MB-231 breast cancer cell line, but not in the MSF-7 or MDA-MB-435 cell lines. This gene and protein provide new markers for distinguishing breast cancer tissue of different types of metastatic potential.

Initially isolated from primary human myeloid leukemia cells, the ufo oncogene (also called Axl or Ark) is a receptor tyrosine kinase (RTK). Its genomic structure is described in Schulz et al., supra., and its differential expression is described in Challier et al., 1996, Leukemia 10:781-787. The ufo protein is a member of a class of RTKs having two fibronectin type III domains and two immunoglobulin-like domains present in the extracellular portion, and is preferentially expressed in monocytes, stromal cells, and some CD34-positive progenitor cells (Neubauer et al., 1997, Leuk Lymphoma 25:91-96). Ufo has an extracellular structure similar to neural cell adhesion molecules, and has direct or indirect binding sites for PLCgamma, GRB2, c-src, and lck (Braunger et al., 1997, Oncogene 14:2619-2631).

<u>eIF-2</u> (SEQ ID NO:71) is a translation initiation factor, and functions as a heterotrimeric GTP-binding protein involved in the recruitment of methionyl-tRNA to the 40 S ribosomal subunit (Gasper *et al.*, 1994, *J. Biol. Chem. 269*:3415-3422). According to the present invention, higher expression is found in two metastatic breast cancer cell lines and not in cell line MCF-7.

eIF-2 is involved in introducing the initiator tRNA into the translation mechanism and performing the first step in the peptide chain elongation cycle. eIF-2 is associated with a 5 subunit molecule having GTP recycling function called eIF-2B (Kyrpides and Woese, 1998, *Proc. Nat'l Acad. Sci. USA 95*:3726-3730, and Kimball *et al.*, 1998, *J. Biol. Chem. 273*:12841-12845).

eIF-2 has subunits alpha and beta. eIF-2alpha is phosphorylated at Ser 51 and then modulates the interaction of eIF-2 and eIF-2B, as described in Kimball et al., 1998, Protein Expr. Purif. 12:415-419, Kimball et al., 1998, J. Biol. Chem. 273:3039-3044, and Pavitt 1998, Genes Dev. 12:514-526. It is reported that by regulating translation initiation, control of cell growth and division in eukaryotic cells is achieved: for example, clotrimazole, a potent anti-proliferative agent in vitro and in vivo, depletes intracellular Ca⁺² stores, which activates PKR, resulting in the phosphorylation of eIF-2alpha, and the ultimate inhibition of protein synthesis and blockage of the cell cycle in G1 phase (Aktas et al., 1998, Proc. Nat'l Acad. Sci. USA 95:8280-8285). Additionally, Kim et al., 1998, Mol. Med. 4:179-190, show that nitric oxide (NO) suppresses protein synthesis in cell types including human ovarian tumor cells by stimulating phosphorylation of eIF-2alpha.

Glutaminyl cyclase (SEQ ID NO:72) is described by Song et al., 1994, J. Mol. Endocrinol. 13:77-86, and is expressed most highly in the most metastatic cell line MDA-MB-435, as compared to less metastatic line MDA-MB-231 and non-metastatic line MCF-7. Glutaminyl cyclase (also called glutamine cyclotransferase) converts glutaminyl-peptides (such as gonadotropin-releasing hormone and thyrotropin-releasing hormone) into pyroglutamyl-peptides, as described in Busby et al., 1987, J. Biol. Chem. 262:8532-8536, Fischer and Spiess, 1987, Proc. Nat'l Acad. Sci. USA 84:3628-3632, and Pohl et al., 1991, Proc. Nat'l Acad. Sci. 88:10059-10063. Cloning and sequence analysis of glutaminyl cyclase derived from a human pituitary cDNA library is described in Song et al., 1994, J. Mol. Endocrinol. 13:77-86. Studies on the catalytic pathway of glutaminyl cyclase and its substrate specificity are described in Gololobov et al., 1996, Biol. Chem. Hoppe Seyler 377:395-398. Assays for the presence of glutaminyl cyclase activity are described in Koger et al., 1989, Method Enzymol. 168:358-365 and Houseknecht et al., 1998, Biotechniques 24:346.

gp130 (SEQ ID NO:73) is transmembrane protein glycoprotein 130. gp130 is a signal transducing shared component of the receptor complexes for the interleukin-6 (IL-6)-type cytokines (Hirano *et al.*, 1997, *Cytokine Growth Factor Rev.* 8:241-252), including IL-6, IL-11, leukemia inhibitor factor (LIF), oncostatin M

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(OSM), ciliary neurotrophic factor and cardiotrophin-1. The N-terminal of gp130 is an extracellular immunoglobulin-like portion of the protein (Hammacher et al., 1998, J. Biol. Chem. 273:22701-22707). Signal transduction including gp130 occurs through the gp130/Jak/STAT pathway 1 (Heinrich 1998, Biochem. J. 334:297-314). The cytokines acting through the pathway that includes gp130 (also called gp130 cytokines) exhibit pleitropic biological activities including immune, hematopoietic, and neural effects (Nakashima and Taga, 1998, Semin Hematol. 35:210-221, Thompson et al., 1998, Neuroscience 84:1247-1255, Hirano, 1998, Int. Rev. Immunol. 16:249-284, Marz et al., 1997, Eur. J. Neurosci. 9:2765-2773, and Betz and Muller, 1998, Int Immunol 10:1175-1184).

gp130 cytokines are reported to control survival and proliferation of myeloma cell lines and primary myeloma cells (Klein, 1998, *Curr. Opin. Hematol.* 5:186-191). gp130 is expressed in the majority of renal cell carcinomas and has an important role in the proliferation of some renal cell carcinoma cell lines (Costes *et al.*, 1997, *J. Clin. Pathol.* 50:835-840).

E-cadherin (SEQ ID NO:75) is a member of a family of glycoproteins responsible for calcium-dependent cell-cell adhesion and is implicated in maintaining cytoskeletal integrity. Epithelial cadherin (E-cadherin) mediated cell adhesion system in cancer cells is inactivated by multiple mechanisms corresponding to the pathological features of the particular tumor type (Hirohashi, 1998, *Am J Pathol 153*:333-339). In general the cadherin system mediates Ca⁺²-dependent homophilic cell-cell adhesion. Transcriptional inactivation of E-cadherin expression occurs frequently in tumor progression, and thus inactivation or downregulation of E-cadherin plays a significant role in multistage carcinogenesis (Hirohashi, 1998, *Am J Pathol 153*:333-339).

E-cadherin is characterized as a tumor suppressor of the metastatic phenotype, as described in MacGrogan and Bookstein, 1997, Semin Cancer Biol 8:11-19, and cadherins are important determinants of tissue morphology including invasive carcinoma as described in van der Linden, 1996, Early Pregnancy 2:5-14, and Yap, 1998, Cancer Invest. 16:252-261.

Mechanisms of action of cadherins are discussed in Daniel and Reynolds, 1997, *Bioessays 19*:883-891. The structure and function of cell adhesion molecules including E-cadherin are described in Joseph-Silverstein and Silverstein, 1998, *Cancer Invest. 16*:176-182, Yap *et al.*, 1997. *Annu. Rev. Cell Dev. Biol. 13*:119-146, and Uemura, 1998, *Cell 93*:1095-1098. Cell adhesion molecules including E-cadherin are potential targets for anti-cancer drugs and therapeutics to treat acute or chronic inflammatory disease as described in Buckley and Simmons, 1997, *Mol Med Today 3*:449-456, Moll and Moll, 1998, *Virchows Arch 432*:487-504.

According to the present invention, E-cadherin is expressed in non-metastatic breast cancer cell line MCF-7, and not in MDA-MB-231 and MDA-MB-435. The expression products are diagnostic markers indicating the metastatic potential of breast cancer tissue samples.

Serpin (SEQ ID NO:76), serine protease inhibitors, are a family of protease inhibitors that inhibit chymotrypsin-like serine proteases (Whisstock et al., 1998, Trends Biochem. Sci. 23:63-67) and that have the unique ability to regulate their activity by changing the conformation of their reactive-center loop; studies of serpin variants provide definition for the functional domains of serpins that control the folding and link serpins mutations to disease (see Stein and Carrell, 1995, Nat. Struct. Biol. 2:96-113). Serine protease cleavage of proteins is essential to a wide variety of biological processes, and the cleavage is primarily regulated by the cleavage inhibitors, as described in Wright, 1996, Bioessays 18:453-464. Members of the serpin family include alpha 1-antitrypsin (AAT) (Carrell et al., 1996, Chest 110:243S-247S), alpha2anti-plasmin (PAI-1 and PAI-2) (Andreasen et al., 1997, Int. J. Cancer 72:1-22), thrombin, urokinase plasminogen activator, and kallikrein (Turgeon and Houenou, 1997. Brain Res Brain Res Rev 25:85-95). Some serpins also have other activities including neuronal differentiating and survival activities (Becerra, 1997, Adv. Exp. Med. Biol. 425:332-237) and tumor suppression (Sager et al., 1997, Adv. Exp. Med. Biol. 425:77-88). PAI-1 and PAI-2 are linked to cancer metastasis, as described in Andreasen et al., 1997, Int. J. Cancer 72:1-22.

pS2 (SEQ ID NO:77) was isolated from MCF7 human breast cancer cells, as described in Takahashi *et al.*, 1990, *FEBS Letters 261*:283-286. pS2 is estrogen-regulated. Speiser *et al.*, 1997, *Anticancer Research 17*:679-684, reported that the pS2 status declined from well to poorly differentiated ovarian cancer. pS2 expression also is associated with a good prognosis in breast cancer patients. According to the present invention, pS2 is expressed in MCF-7 cells, but not in two metastatic breast cancer cell lines

pS2 (presenilin-2 or trefoil factor 1 (TFF 1)) is a trefoil polypeptide normally expressed in the mucosa of the gastrointestinal tract, and found ectopically in gastrointestinal inflammatory disorders and various carcinomas (May and Westley, 1997, *J. Pathol. 183*:4-7. pS2 is expressed in breast cancers (Poulsom *et al.*, 1997, *J. Pathol. 183*:30-38). pS2 is a pleitropic factor involved in mucin polymerization, cell motility (Modlin and Poulsom, 1997, *J. Clin. Gastroenterol 25*(1):S94-S100), cell proliferation and/or differentiation, and possibly in the nervous system (see Ribieras *et al.*, 1998, *Biochim. Biophys. Acta. 1378*:F61-F77).

LIV-1 (SEQ ID NO:78) is an estrogen-regulated protein reported in the MCF-7 cell line (Green *et al.*, GeneBank submission Accession No. U41060). According to the present invention, LIV-1 is expressed in MCF-7 cells, but not in two metastatic breast cancer cell lines.

Leucine-isoleucine-valine -1 (LIV-1) and other members of the LIV family (LIV-2, 3, and 4) are binding proteins that represent a transport system for branched chain amino acids in *E. coli* as described in Yamamoto *et al.*, 1979, *J. Bacteriol. 138*:24-32, and Yamamoto and Anraku, 1980, *J. Bacteriol. 144*:36-44. A human homologue to LIV-1 is both estrogen and growth factor inducible in MCF-7 human breast cancer cell line (El-Tanani and Green, 1997, *J. Steroid. Biochem. Mol. Biol 60*:269-276; El-Tanani and Green, 1996, *Mol Cell Endocrinol 124*:71-77; and El-Tanani and Green. 1996, *Mol Cell Endocrinol 121*:29-35).

GTP-binding protein (SEQ ID NO:79) is a member of the family of guanine nucleotide-binding regulatory proteins, G proteins. The protein is expressed in MCF-7 cells, but not in two metastatic breast cancer cell lines.

G proteins provide signaling mechanisms for the serpentine family of receptors as described in Dhanasekaran and Prasad, 1998, *Biol. Signals Recept 7*:109-117. Studies indicate that the alpha as well as the beta gamma subunits of the GTP-binding proteins are involved in the regulation of several cellular responses, some of which responses are critical to the regulation of cell growth and differentiation (Dhanasekaran and Prasad, 1998, *Biol Signals Recept 7*:109-117). G protein coupled receptors regulate the mitogen activated protein kinase pathway as described in Russell and Hoeffler, 1996, *J. Invest. Dermatol Symp Proc 1*:119-122, and thus play a role in controlling cell growth. GTP binding proteins are also implicated in the regulation of intracellular transport as described in Ktistakis, 1998, *Bioessays 20*:495-504.

Chemokines induce various intracellular signaling pathways in natural killer cells by activating members of GTP binding proteins as described in Maghazachi and Al-Auokaty, 1998, FASEB J. 12:913-924. Heterotrimeric GTP binding proteins regulate distinct signaling pathways, some of which in turn regulate the activity of Na+/H+ exchanger proteins as described in Voyno-Yasenetskaya, 1998, Biol Signals Recept 7:118-124.

<u>Desmoplakin</u> (SEQ ID NO:84) is a member of a family of proteins that serve as cell surface attachment sites for cytophasmic intermediate filaments.

<u>Vimentin</u> (SEQ ID NO: 80) is a member of the intermediate filament gene family (Evans, 1998, *Bioessays 20*:79-86. Intermediate filaments are a major component of the cytoskeleton of higher eukaryotes. Vimentin gene knockout mice indicate degeneration of the cerebellar Purkinje cells (Galou *et al.*, 1997, *Biol Cell 89*:85-97). Vimentin is positive in immunohistochemical reactions of sarcomas and related lesions (Gaudin *et al.*, 1998, *Am J Surg Pathol 22*:148-162), and of desmoplastic small round-cell tumors and their variants (Gerald *et al.*, 1998, *J. Clin. Oncol. 16*:3028-3036). Vimentin is also expressed in neoplasms showing follicular dendritic cell differentiation as described in Perez-Ordonez and Rosai, 1998, *Semin. Diagn. Pathol. 15*:144-154, and in biphasic carcinomatous-sarcomatous malignant mixed mullerian tumors as described in Guarino *et al.*, 1998, *Tumori 84*:391-397.

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Cytochrome C Oxidase (CcO) (SEQ ID NO: 81) is the terminal enzyme of the respiratory chain of mitochondria and aerobic bacteria: it catalyzes electron transfer from cytochrome C to molecular oxygen, reducing the oxygen to water (Michel et al., 1998. Annu Rev Biophys Biomol Struct 27:329-356). Cytochrome C oxidase is a member of the superfamily of quinol and cytochrome C oxidase complexes that are related by a homologous subunit containing six positionally conserved histidines that ligate a low-spin heme and a heme -copper dioxygen activating and reduction center as described in Musser and Chan, 1998, J. Mol. Evol. 46:508-520. Cytochrome C and ubiquinol oxidases are membrane-bound redox-driven proton pumps which couple an electron current to a proton current across the membrane (see Karpefors et al., 1998, Biochim Biophys Acta 1365:159-169). Analysis of mutant forms of cytochrome C oxidase is described in Mills and Ferguson-Miller, 1998, Biochim Biophys Acta 365:46-52. Nitric oxide inhibits respiration at cytochrome C oxidase, as described in Torres et al., 1998, J. Bioenerg Biomembr 30:63-69.

Heat shock protein 90 (hsp90) (SEQ ID NO: 82) acts as a chaperone molecule in association with the glucocorticoid and progesterone nuclear receptors, and has A, B, and Z regions for facilitating these interactions (Dao-Phan et al., 1997, Mol Endocrinol 11:962-972). Levels of hsp90 are reported elevated in active systemic lupus erythematosus (Stephanou et al., 1997. Biochem J 321:103-106). Increased hsp90 expression is implicated in regulation of forms of cell injury that lead to programmed cell death as described in Galea-Lauri et al., 1996, J. Immunol. 157:4109-4118. Hsp90 is upregulated in regenerating fibers and diseased fibers of Duchenne muscular dystrophy (Bornman et al., 1996, Muscle Nerve 19:574-580), and is a candidate substrate for proteolysis during ionizing radiation-induced apoptosis of some breast cancer cells (Prasad et al., 1998, Int. J. Oncol 13:757-764). Hsp90 is involved in dislocation of the mutant insulin receptors from the endoplasmic reticulum to the cytosol as described in Imamura et al., 1998. J. Biol. Chem. 273:11183-11188. and associates with and activates endothelial nitric oxide synthase as described in Garcia-Cardena et al., 1998, Nature 392:821-824.

Integrin alpha 6 (SEQ ID NO: 83) is in the family of integrins. heterodimeric, cation dependent cell membrane adhesion molecules that mediate cell-cell and cell-matrix interactions. Integrin alpha 6 is a component of the hemidesmosome complex (Jones et al., 1998, Bioessays 20:488-494). Integrins maintain tissue integrity and regulate cell proliferation, growth, differentiation, and migration. (See Thomas et al., 1997, Oral Oncol 33:381-388). In oral squamous cell carcinomas there is a variable loss or reduced expression of integrin alpha 6, as described in Thomas et al., 1997, Oral Oncol. 33:381-388. Alpha 6 integrin also plays an active role in invasion of intestinal and diffuse-type cells of representative gastric carcinoma cell lines as described in Koike et al., 1997, J. Cancer. Res. Clin. Oncol. 123L:310-316.

Osteogenic protein-1 (OP-1) (also called BMP-7) (SEQ ID NO: 85) is a morphogenetic factor (and a member of the bone morphogenetic protein (BMP) family of growth factors) and is highly expressed in kidney and involved in tissue repair and development (see Almanzar et al., 1998, J. Am. Soc. Nephrol. 9:1456-1463). OP-1 is also expressed in the developing nervous system and can induce dendritic growth in sympathetic neurons as described in Guo et al., 1998, Neurosci. Lett 245:131-134. OP-1 stimulates cartilage formation as described in Klein-Nulend et al., 1998, J. Biomed. Mater. Res. 40:614-620.

OP-1 induces down-regulation of insulin-like growth factor binding proteins (particularly IGFBP-5) thus affecting IGF-1 in the context of bone cell differentiation and mineralized bone nodule formation as described in Yeh *et al.*, 1997, *Endocrinology 138*:4181-4190. OP-1 can be used as a bone graft substitute to promote spinal fusion and to aid in the incorporation of metal implants (Cook and Rueger, 1996, *Clin. Orthop. 324*:29-38). The three dimensional structure of OP-1 is reported in Griffith *et al.*, 1996, *Proc Nat'l Acad Sci 93*:878-883.

The protein encoded by SEQ ID NO:56 is a putative secreted protein and is highly expressed in fat tissue.

Table 1. Novel Differentially Expressed Metastatic Marker Polynucleotides

TRANSCRIPT NUMBER	SEQ ID NO:	non- metastatic breast MCF-7	breast cancer cancer metastatic to bone and/or lung MDA-MB-MB-435		low metastatic from colon KM12C	high metastatic from colon KM12L4A
901	1	_	+	-		
907	2	-	-	+		
9102ь	3	+	-			
9114	4	-	-	+		
9121a	5	-	+	<u>-</u> ·		
9129	6	+	-	+		
9139a	7	+	-	-		
9143b	8	+	-	-		
9157Ь	9	-	-	+		
9166	10	+	•	-		
9170ь	11	-	+	-		
9190a	12	+				
9191	13	-	-	+		
9216	14	-	-	+		
9224c	15	+	-	*		
9230ь	16	+ :	-	-		
924	17	+	-	-		
9242a	18	-	+	•		
9259a	19	-	-	+		
9261	20	-	+	-		
9272	21	+	-	-		
9293ь	22	-	+	-		
9304b	23	+	-	-		
9307a	24	-	+	-		
931	25	+	-	•		
9313	26		-	+		

TRANSCRIPT NUMBER	SEQ ID NO:	non- metastatic breast MCF-7	breast cancer metastatic to bone and/or lung MDA-MB- 23 l	breast cancer metastatic to lung MDA- MB-435	low metastatic from colon KM12C	high metastatic from colon KM12L4A
9316	27	+	+	-		
9318b	28	+	-	•		
9320a	29	-	-	+		
9330b	30	-	+	-		
9335	31	+		-		
9337	32	+	-	+		
9342b	33	-	+	-		
9343c	34	+	-	-		
9350e	35	-	+	-		
9351b	36		+	-		
9361	37	+				
9368	38	-	+	-		
9373b	39	-	-	+		
9385a	40	-	- +			
9386c	41	-	-	+		
9388d	42	+	-	-		
9390	43	+	-	-		
9393	44	+	-	-		
9396	45	-	+			
944b	46	+	-	-		
951	47	+	-	-		
953	48	-	-	+		
954a	49	+	-	-		
968	50	+	-	-		
971	51	+	-	-		
983c	52	-	+	-		
985	53	+	-	-		
990	54	+	-	+		

4	9

TRANSCRIPT NUMBER	SEQ ID NO:	non- metastatic breast MCF-7	breast cancer metastatic to bone and/or lung MDA-MB- 23 l	breast cancer metastatic to lung MDA- MB-435	low metastatic from colon KM12C	high metastatic from colon KM12L4A
998	55	-	-	+		
316	56	+	-	-	+	-
126c	57	-	-	+		
207-4	58	-	+			
265-3	59	+	-	•		
29B	60	-	-	+		
305B-25	61	+	-	<u>-</u>		
326B-39	62	+	-	-		
34B-11	63	-	-	+		

- + indicates differential expression as identified in differential display
- indicates absence in differential display

For transcript number 316, reverse transcription PCR (RT-PCR) was used to detect expression in the breast cancer cell lines.

Table 2. Differentially Expressed Metastatic Marker Polynucleotides

TRANSCRIPT NUMBER	protein	SEQ ID NO:	non- metastatic breast MCF-7	breast cancer metastatic to bone and/or lung MDA-MB- 23 I	breast cancer metastatic to lung MDA-MB- 435
902	osteopontin	64	-	-	+
9112	nip	65	-	+	-
9132	Ca-dependent protease	66		+	-
9158	IGF-R	67	+	-	-
9174	ILGF-BP5	68	÷	-	-

TRANSCRIPT NUMBER	protein	SEQ ID NO:	non- metastatic breast MCF-7	breast cancer metastatic to bone and/or lung MDA-MB- 23 l	breast cancer metastatic to lung MDA-MB- 435
9177	lactate dehydrogenase	69	· <u>-</u>	+	+
9202	ufo TKR	70	-	+	-
9210	elF2	71	-	+	+
9212	glutaminyl cyclase	72	-	-	+
9213	gp130	73	-	-	+
9222	TGFb-II	74	-	+	-
9232	E-cadherin	75	+	-	-
9239	serpin	76	-	+	-
9250	secreted pS2	77	+	-	-
9260	LIV-1	78	+	-	<u>-</u>
9315	GTP-binding protein	79	+	-	-
9317	vimentin	80	-	+	-
938	cytochrome C oxidase	81	+	-	-
9382	Hsp 90	82	-	-	+
9394	integrin a6	83	-	-	+
956	desmoplakin	84	+	-	-
970	osteogenic protein	85	+	-	-

- + indicates differential expression as identified in differential display
- indicates absence in differential display

Within the scope of the invention are variants of the proteins described above. A variant is a protein encoded by a polynucleotide wherein the global sequence identity of the DNA, as compared to the corresponding SEQ ID NO: herein, is at least 65% as determined by the Smith-Waterman homology search algorithm as implemented

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in MPSRCH program (Oxford Molecular) using an affine gap search with the following search parameters: gap open penalty of 12, and gap extension penalty of 1. The protein encoded by the DNA having the sequence identity described above will exhibit the percent activity described in the preceding paragraph.

Also within the scope of the invention are fusion proteins comprising the proteins and variants disclosed herein. Proteins preferably used in fusion protein construction include beta-galactosidase, beta-glucuronidase, green fluorescent protein (GFP), autofluorescent proteins including blue fluorescent protein (BFP), glutathione-Stransferase (GST), luciferase, horse radish peroxidase (HRP) and chloramphenicol acetyltransferase (CAT). Additionally, epitope tags are used in fusion protein constructions, including Histidine (His) tags, FLAG tags, influenza hemagglutinin (HA) tags, Myc tags, VSV-G tags, and thioredoxin (Trx) tags. Other fusion constructions can include maltose binding protein (MBP), S-tag, Lex A DNA binding domain (DBD) fusions, GAL4 DNA binding domain fusions, and Herpes simplex virus (HSV) BP16 protein fusions.

These fusions can be made by standard procedures in the art of molecular biology, and many are available as kits from, for example, Promega Corporation (Madison, WI); Stratagene (La Jolla, CA); Clontech (Mountainview, CA); Santa Cruz Biotechnology (Santa Cruz, CA); MBL International Corporation (MIC, Watertown, MA); and Quantum Biotechnologies (Montreal, Canada).

The proteins of the invention, and variants as described herein, can also be used to detect protein interactions in vivo, using the yeast two-hybrid system, for example as described in U.S. Patent No. 5,674,739.

In addition to the ribozyme and antisense constructs described above, the polynucleotides of the invention can be used for inhibiting transcription via triple helix formation as disclosed in U.S. Patent No. 5.674,739.

Those skilled in the art will recognize, or be able to ascertain, using not more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such specific embodiments and equivalents are intended to be encompassed by the following claims.

All patents, published patent applications, and publications cited herein are incorporated by reference as if set forth fully herein.

CLAIMS

We claim:

- 1. An isolated and purified human protein comprising an amino acid sequence which is at least 85% identical to an amino acid sequence encoded by a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.
- 2. The isolated and purified human protein of claim 1 wherein the amino acid sequence is at least 95% identical.
- 3. The isolated and purified human protein of claim 1 wherein the amino acid sequence is encoded by a sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.
- 4. A fusion protein which comprises a first protein segment and a second protein segment fused to each other by means of a peptide bond, wherein the first protein segment consists of at least six contiguous amino acids selected from an amino acid sequence encoded by a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.
- 5. A preparation of antibodies which specifically bind to a human protein which comprises an amino acid sequence encoded by a nucleotide sequence selected from the group consisting of SEQ ID NOS:1-63 or the complement thereof.
- 6. A method for detecting metastatic tumor cells in a tissue sample, comprising the step of:

measuring in said tissue sample an expression product of a gene which comprises a coding sequence selected from the group consisting of SEQ ID NOS:1, 2, 4, 5, 9, 11, 13, 14, 18, 19, 20, 22, 24, 26, 29, 30, 33, 35, 36, 38-41, 45, 48, 52, 55, 57, 58, 60, 63-

66, 69-74, 76, 80, 82, and 83, wherein a tissue sample which expresses the product is categorized as containing metastatic tumor cells.

- 7. The method of claim 6 wherein the expression product is protein.
- 8. The method of claim 7 wherein the protein is measured using an antibody which specifically binds to the protein.
- 9. A method for detecting metastatic tumor cells in a tissue sample, comprising the step of:

measuring in a tissue sample an expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:3, 7, 8, 10, 12, 15-17, 21, 23, 25, 28, 31, 34, 37, 42-44, 46, 47, 49-51, 53, 59, 61, 62, 67, 68, 75, 77-79, 81, 84, and 85, wherein a tissue sample which does not express the product is categorized as metastatic.

- 10. The method of claim 9 wherein the expression product is protein.
- 11. The method of claim 10 wherein the protein is measured using an antibody which specifically binds to the protein.
- 12. A method for determining metastatic potential in a tissue sample, comprising the step of:

measuring in a tissue sample an expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:1, 2, 4, 5, 9, 11, 13, 14, 18, 19, 20, 22, 24, 26, 29, 30, 33, 35, 36, 38-41, 45, 48, 52, 55, 57, 58, 60, 63-66, 69-74, 76, 80, 82, and 83, wherein a tissue sample which expresses the product is categorized as having metastatic potential.

13. The method of claim 12 wherein the expression product is protein.

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14. The method of claim 13 wherein the protein is measured using an antibody which specifically binds to the protein.

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15. A method for determining metastatic potential in a tissue sample, comprising the step of:

measuring in a tissue sample an expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:3, 7, 8, 10, 12, 15-17, 21, 23, 28, 31, 34, 37, 42-44, 46, 47, 49-51, 53, 59, 61, 62, 67, 68, 75, 77-79, 81, 84, and 85, wherein a tissue sample which does not express the product is categorized as having metastatic potential.

- 16. The method of claim 15 wherein the expression product is protein.
- 17. The method of claim 16 wherein the protein is measured using an antibody which specifically binds to the protein.
- 18. A method of predicting the propensity for metastatic spread of a breast tumor preferentially to bone or lung, comprising the steps of:

measuring in a breast tumor sample an expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NO:1, 5, 11, 18, 20, 22, 24, 30, 33, 35, 36, 38, 45, 52, 58, 65, 66, 70, 74, 76, and 80,

wherein a breast tumor sample which expresses the product is categorized as having a propensity to metastasize to bone or lung.

19. A method of predicting propensity for metastatic spread of a breast tumor preferentially to lung, comprising the steps of:

measuring in a breast tumor sample an expression product of a gene which comprises a sequence selected from the group consisting of SEQ ID NOS:2, 4, 9, 13, 14, 19, 26, 29, 39-41, 48, 55, 57, 60, 63, 64, 72, 73, 82, and 83,

wherein a breast tumor sample which expresses the product is characterized as having a propensity to metastasize to lung.

20. A method of predicting propensity for metastatic spread of a colon tumor, comprising the steps of:

measuring in a colon tumor sample an expression product of a gene which comprises the nucleotide sequence shown in SEQ ID NO:56,

wherein a colon tumor sample which expresses the product is characterized as having a low propensity to metastasize.

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aanatnatac ctantaanga acnctgtaca ntgccncaag cangtganga ccncccacga
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gtttacatna atacaatnot gaaacnacno aggotggttt tatatotaca tatttgactt
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accactaten cantaaagtt tngcacettt eneegaacga aaanaacece eentnntgnn
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ttcttttnaa aanacentng nneenenttn eegtenenee eennatantn nnennateee
cccctctncc nntccntnnn cgtaannggc gtngcttntg cngtntntgt cccgttttcc
                                                                        420
```

```
tecgettngt entttnteta tatnggetnn tnttatneen ngecettegt encetnnngn
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ttcqtctqtn cntaqtcctc ntnctngagc cccanttgnt acttcnngct tcnnctccgc
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atteentete egenennane nennntetea nannatgnne nntnnetnen neenatnene
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cctnanagnt tcgnctagac cntcnacntt gtntcccgnn ctcttagngn tctgctncta
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qtqtntnnct catctcctct ncttctctct cctttgacnc ngnnenctcc atcntnntct
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quettetea tenennung eecetneten ennagtnign gigenennne tinnuntena
                                                                       780
nctngtegec teegtttten actnnnncen nngengnneg nnngetettt etntenntta
                                                                       840
gactnacctt ntctgnnnnn tcannctagc nctgtccntc tctnntctgc atcnttanac
                                                                       900
atettnnten ecenetegea nentnetntt nachetenea taegttneen nneteagtee
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qcaqnnnnqt thenthengt entetegegn etennnteet etethnnaen encetggtet
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negneteget conneceatn enthectegt tgntennnnt ennataegtn theangeene
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ntctctccnc tn
                                                                      1092
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      <212> DNA
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      <221> misc_feature
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taatgccaag ggagaanagc cagtacacta tatggtttat actctttatc cctttattca
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cttttgtgca ccctggtgca catctgactg ttgtcctanc canaaactct ctgaggccac
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tgaaagaaca gtggccctat cgatttcatt cctaggtctc aaaaatacna tgtngccttg
                                                                       420
taacataatt agggacagca cctctatttc acaattataa tctaaggtag gataagacga
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cacagcagca ataaacttac aagt
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      <221> misc_feature
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atgengeeaa gettntttat tgaaaantee taattntatt gneegtntag taacatgttt
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gttcnacaan gctaatttct nataaancaa aacacannnt tttcttataa gtngtataaa
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ttatttnatt tacagaaact tgtttcaaaa canatgnact anntatttct nctcttttaa
                                                                       300
atanccanac taattttcta tccctngaca tctgttcatg ttctatncag cagccaacac
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aaagtccanc tgagagctct tgattaangt gtncgnatta tctagctact tccnacgttt
                                                                       420
tnggngcnng aaatgncttt taanancctg gcctcaaaaa anaaaaanan cccccgnnn
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aggggnnttc cntntanaaa aanggntcnc tcnnccngtn ngagactgtc tccctgnntn
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ngnnnntege thinateang ngeenenang etencenten etnnngeatt ngathnntan
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cnnnctgaga tgngnntang ctgntncntn ngtgtcntan gtctcgacgt tgnntggntn
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tangnancgn cnntntnnnc nnattgncga gngnntaagt gtgctcttct cntnacntct
                                                                       720
ntennnanen tetnngatgt tnataeggee gtgettnett atenntgana negntetnan
                                                                       780
nanntnegna tgagnntnta etgenenent gtgteatett tetetetant gtgtnetnna
                                                                       840
nncnngtnat tncgcnnnac tgntantnag tggtatnnag anntcgnncg cnngngccnn
                                                                       900
                                                                       960
tttnnctgtn gnnatnagnt ntcanganat tnatncnntc tncgtgatag anagntnagt
                                                                      1020
qnnqqntctg actgatncgt gtcctagtnn cngtgacatc gnncgttann gtcngcactc
                                                                      1080
tagtanannt nagtnngang ntgtanatnn ntctcntgtt tcagtnnagn cccncgagcg
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                                                                      1132
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      <212> DNA
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      <221> misc feature
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antnagnaat gcctttttnc tgagggcntt nggnnntcat nnangggngt gnggnggntt
                                                                       180
ncacctgtaa taccaccact ttncnatgcc actgccngtg natcaccngn ngtaaggact
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tcaanaccaq ccttatnaac ntgggnaaac cntntntcta ctaaaaatnc tnnaantatc
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tgngcnnngt ngngcgttct tntannnccn gctgnacnng angncngngn angntantcg
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cntgaacntg ncntgttana gtngcantga gcctaaatca cantgatgta ttnncatctg
                                                                       480
ggacgacacg ancngacgac tenegtaeth aaaaaaaaaa necenttnng ggggggtttt
tnnnggtatt anntatantt ggagaanttt gggtcannng aatattntta catgaaaaat
                                                                       540
naggaataac thtathtgtg tacattgggt thnaaanang acantantgg nhctaaacth
                                                                        600
                                                                        660
ttngggngg aggggnnatt agggnnttaa ttnggnnnct tnnaaanncn nntnnngtat
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nanaanantn tttnnanaag ngnantngnt ttaaancctn aangnttnnn tnctnttann
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ttnnaannnn anannn
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      <211> 690
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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      <223> n = A, T, C \text{ or } G
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caqtattact qttttgctaa gccgcttcat tcatgcctac acaatttttt tttaaaaggg
aactttagtt aattaagtga taagggactt aaatatgaat tanaatggtg cagaaagaga
                                                                        180
taccttttct ggatatttta aagtttaaag gtcantttct cttaatctga ttatgtgcac
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atatgaaaat ggcacatcat atacatgtaa aatcaggcag tatncattta ttaattactg
                                                                        300
tatttgacaa aggaaactct taaattataa tgtgaaacct ggttttatga aaccaatgac
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taqtqcanca tttcagcata tgcaaaaaaa aaanncctnt tggngngctg tttacaaagg
                                                                        420
aaattgttgg atttcacgat ggtttcagga naanaaggtt ttcntcatcn agggtaaacn
                                                                        480
tcccggataa ggcntngntt taatntnntt annccnnccn atngntaann gtggaaatta
                                                                        540
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```
ancetetgaa naaaanance cacntnnttn geettggget tnantetntt tggengnane
                                                                         600
naaaggnnct tnccaggtnt cntgnngggc cngnngaann ataannaann nggggnnctt
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      <211> 395
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      <221> misc feature
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                                                                         120
cagaggttna aattatcaga cagaaccttt aanaataatt atgattaatg tgttaaaatt
                                                                         180
ctagtggaaa agataaataa catgctcagg anattttagc anagagatag aaactatntn
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ngaagctcaa atgaaaatgc taggaaatga aaagcagtat tggaggtgaa agattccttt
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      <220>
      <221> misc_feature
      <222> (1)...(331)
      \langle 223 \rangle n = A,T,C or G
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                                                                         120
ntcgctacag gagggaacgt gaaaagaaca tctccagagg aactggtgaa tgaccacgcc
                                                                         180
cqagagaaca gaatcaaccc cgaccaaatg gaggaggagg aattcataga aataacgact
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gaaagaccta aaaagtagca agaagctaca tccctcaaac ttcggcaatg aaaataaagt
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ttgagaagct caaaaaaaa aanccctttt g
                                                                        331
      <210> 12
      <211> 693
      <212> DNA
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      <220>
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      <223> n = A,T,C or G
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caaactgacc ttactttctt gaagacggaa ttgtagtatg gtcgagctca tgctttttgt
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agtaggccat ncaaattcga ttgactggct aaaaaaagatt gttagtggag gctggaagaa
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acattttqqc tqatqataqa tgaatagagc ttggaacaat caaaaggaaa agcagaaagt
ctatacctat tcataagaaa aagttagtat gtttaccgaa cattatnaaa gaattatgac
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attttcaaag ttttaaaatt ttattttgta gggacggggt ctcattgtgt agcccacnct
ggtctgtttc ttgaggattt actatanact gggctgtatt caaagcattg gggatacagg
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catqaatqag cccccattgc ctgaacttac cattcaatct gggcagtgaa agaanaggga
tgntgggaga nccttacaaa gatgaaatgt cgctaactgg agaaatccct actttcagtc
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agactgaann ggaacaggta gtnactgtgg gtagccctct ttgggnangg gtngattttc
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cacatgtgcc cagttaaggg ccnagaacat taa
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      <211> 305
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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      <223> n = A, T, C \text{ or } G
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tccttggnat canannatat cntggccnaa gaaccncnca ccntctntgg gttagaaata
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ccgctntatn gngtatgagg ggatngggcn tacgnnataa tttnctatng ganggtattn
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ccgcactant gacnagttct ttctnnggtc catttnnaac nacantnttg acattgntga
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tctgcaannc tgtaaaatag tcttncagtg ggcaatnnnt gcacaactgg gttnggtntc
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                                                                        305
anaca
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      <211> 308
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(308)
      <223> n = A, T, C \text{ or } G
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ggcgctctag atatantgcc ccaaaggaaa gagnacaaag tnttccnccc ntagttctac
                                                                        120
natgnetate enetateace thetgntten naagntttht aaaaataaat tetettgtat
                                                                        180
ancatecnat ateneacegg tecaaagege aacaatetge aatteanaan ttecaacaat
                                                                        240
cnatntatgn actttentag gtccggtgtt ctaanatnta atattetaac acttactete
                                                                        300
                                                                        308
agatetta
      <210> 15
       <211> 304
       <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
       <222> (1)...(304)
       <223> n = A,T,C or G
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ngtnaaggga tatttattcc tgttttaaaa ggatacaacc aaggtaggga aggcttcgtt
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attggtgatt attcagaaga cctattttct ttacatatgc tatggaaaca atactqtttt
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ccgctacaga atacagttta tgattatact tttgtaaatt gcctgctttt cccctgtcat
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ctgctaattc caatttgata ctgttctgtg ttcaaaaata cagcatgagc aagctgtaat
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ggtgcctgtc gagagtccca gctgcttggg gggctaaggt gggaggatca tttgagccca
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ggag
                                                                        304
      <210> 16
      <211> 703
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(703)
      <223> n = A, T, C \text{ or } G
      <400> 16
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                                                                         60
gattattatt ttagatccta catatacttt tatcagtaga atqatttcat tnaqatqtat
                                                                        120
aatgaaaaag ggtaatgcaa aaattatgta atagatacca aattagggaa gtttggcaat
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ttcaatggca tatttttagt caaggnacac agatggcagt gccataagca agtctataaa
                                                                        240
tatcggctgc agccatcccc ctcattttaa atgttgccct aataatcaat gcagttaaca
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agtatattgg ctgtgtca tgaaatagtt catgttcaga tggaaatgtt aggttactgt
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tttnnnangn acngggctgg attcaaanca ttggggatnc angnttnaat gngnccccat
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tttnnangnt nnaantgtnn ncttactggn gnaaannncc ntaanntttn nnnntnnnn
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ngnaangggg naannnnnn ntnancttnt gggggagnen nttntggggn anggggggnt
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nnttnnnncn tnnnggccnn nnnnggggcn nnaaantttt tgn
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      <211> 171
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
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      \langle 223 \rangle n = A,T,C or G
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taatanaatc ttcaacatgg cnatccacnc tattccaata atgaaatgca aatttccctq
                                                                        120
ccttctttac tanggtcatt tntagattct tgaggaatga gttctactct t
                                                                        171
      <210> 18
      <211> 689
      <212> DNA
      <213> Homo sapien
      <221> misc_feature
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<222> (1)...(689)
      <223> n = A, T, C or G
      <400> 18
antnngcttn ggtactaagc agaatcactt ncttgggaac tccatgtaac tngtggcttt
                                                                         60
tgtgattgaa atagcatcag taaangtctg accctgtggt aaagacacat atgngcgtgg
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accnggctat gtctgacttt gtgctgctca ggacactctc tgtnaccaaa agngagagan
                                                                        180
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cctqqannac ctcangggt canatgtttg aaggagctgc tgagtatcct ggcaggcanc
anagcettae cateagtttg etgeatggaa ggetgtgtge etetatttee etgetatttg
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ctgagagggc tggggagatg ttngaaggaa agcttggctg gggagctgaa tctggcctgt
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ggtacatgct tggtaactgg tggccaggan acccgggngt gtgtnctggg actgtcncac
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tctqctqacc aqqqtattga aagtccccnc tcaaanacac agaatntntc tgaccaaggg
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tangtatgan atgachtgtg gagcactttg nataaactgg ttctcatngg nggtcccctt
                                                                        600
qaanaqqtqc tnnatctgtt caaaaatacg tggctgagct ntanacccng natcctctgt
                                                                        660
                                                                        689
cagagacatg ggcaggggga ctcaatgct
      <210> 19
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      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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tgctcattca acagtcttgc attcagtagg tgtttgacat cacctactat gtgncaggct
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ctatgctang nactggggat acaggagaga ntnaagcgta aagtctttgg tctcaaggaa
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tataacqaaa tataaagggt ttgggagcaa aaaanaaacc cnnttgtggg gntctntncc
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nctctgatga agettactta ettttaacet tneettetee tttaaaggtg ttteetggtt
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cccctttcct ttacagattg gttattggtc ttgctgagga gtaggactac aattnccagc
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                                                                        660
cccntaatgg taaaatngta ttaaaangtg gacctttgac aaataaattg nttcgatttc
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С
      <210> 20
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      <212> DNA
      <213> Homo sapien
      <220>
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```
ncatchtgan cannatcccc catchnccat atgntgatha nnacaaacca thctattncq
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aatcggta
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      <211> 298
      <212> DNA
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      <220>
      <221> misc feature
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      <223> n = A,T,C or G
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tachanggac theghnanet gggatenggg httachttgt teathgthag agtghnanen
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aagtanatgn taggnataaa gatgttncgg gagatgggtc tacaaantct tttnaagatg
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ntcatcttga anannatcaa gtgtgnttgg tataatgact atcattatac aatgtcaa
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      <211> 591
      <212> DNA
      <213> Homo sapien
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      \langle 223 \rangle n = A,T,C or G
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ctgaagacat tgaataaccc tgggcagtgg ttcttaggca gatactctag atqctttatq
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gacaatatta ttttcattgg atgattctgg agctctatta ggagaaaagt aatcatttta
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ggtcttaaag acttcaagaa aatacaggtt atcaatttat tttaaatctc attgtttcca
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ctaggtcaga aggaaacata ccactctcat ggttcatagt attcactgta tgtatgctag
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ggaaaagact tgctccagtc tcctcctcag ttctgtgcct gagaaccact gctgcatata
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tttgttttta aattttgtat tgaactgtta attgaagctt taaaaqcata tatqaaatqt
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      <210> 23
      <211> 755
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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      <223> n = A, T, C \text{ or } G
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                                                                         60
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120
ttcaaqaqat taggagactt gttcaaagac acacagctgg taagtgatgg aggcaggatt
taaacctggg tttcactgca tttcccatca ctggctttta gccatgatgc tctactgtgt
                                                                       180
                                                                       240
aaccetetta attettgace tgtggetata aagtatgtat tgagagacag geceteeetg
agataacttt ccagccttga caaaggcaca cccttggttc attccttgga gtgtaggacc
                                                                       300
                                                                       360
taqattqtqa caagcccaga tgagtgtgtc tggcagaggg gagcagatct gaggccacca
tatgtgttca cctagcccta aggagtgcca gcttcgctgg tatttgtaca gcttccatca
                                                                       420
ggactgctca ttggccacgt tctttcctct ccctgccacg ttgattaata ctcacataaa
                                                                       480
ttaatgctca cattagtgtt caagtatgca aatgagtgct taaaatcatc actcacacaa
                                                                       540
                                                                       600
tgaccagact gaggatataa cacacaagag ccctttctt ggtaacccca caatcatgca
gatgtgttga cttctctgca ttaccagtct ggtaggcagg gggatatgac agttagaaac
                                                                       660
                                                                       720
aqtctttcan acagcagttc tcaacaccag gtcccttgct gcacaatcga atcacctggg
                                                                       755
ggtttaaaaa aatatcatgc cagtcagcca cnntt
      <210> 24
      <211> 513
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1) ... (513)
      <223> n = A,T,C \text{ or } G
      <400> 24
ctttctaccc aacaagcata gaatatacat tgtatacatc agaaacacgg gacattctcc
                                                                        60
aaaatagacc atatgatagg gcacaaaaca agtctcagta aatttaagaa aatcagaatt
                                                                       120
atatcaagta ctctctcaga ccacagtgga ataaaattgg aaattaattc cgaaaggaac
                                                                       180
                                                                       240
actcaaaaqc atgcaaatac atggtaatta aataacctac tcctgaatga ttgttgggtc
nacaatgata tcaagaggga aatttaaaaa ttctttgaac tgaacgataa tagtgacaca
                                                                       300
qcctatcaaa aactctggga tacagcaaaa gtggaggtaa gaagaaaatt catagcatta
                                                                       360
aatgcctata tcaaaaatct gaaagagcac aaataaacaa tctaaggtca ccctcncaga
                                                                       420
                                                                       480
attqqaqaaa ctagaacagt ccaaatccaa accongcaga agaaaagaaa taaccaaatc
                                                                       513
cgaacaaaac taaatgaatt gaaaaaaatc ccc
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      <211> 574
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(574)
      <223> n = A,T,C or G
      <400> 25
cgatccaaga gattagaanc ccntggagtg gagcatgctt cnctanaatn ccacctgatn
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cttggctnaa nacantnngc tctantttgc tttgtgcccg tccacacaan ctaaaaacaa
                                                                        120
gggatggggg gaccncnagt gtctaatatn cntaatatcc ntccncnggc aaatgaatac
                                                                        180
tttttacaca cttgtanntt ntggagggan ggggtnatna tgaggggaan gggaaaggat
                                                                        240
                                                                        300
qaqqaqaaat ccaggatnan angtototto gtoototona gactnootoa cactotntgt
                                                                        360
ggtnaccngg gttcgttntg tccaatggca gacattatac tccatantct acccnggctt
                                                                        420
nntcqqqttq qqacqccann actcccccna gtngtnnccc ccnancagcn atacacaagt
                                                                        480
ntqaacqqgt tttgtggcca ntcatcgcaa tgaccttntc ctcnactcna agaaaantaa
acceptace congattggt ttctaaatct ttcaccccat ctaaaataga aagchetnag
                                                                        540
tgggangggt tnatcccccc nttaccntta aaac
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<210> 26
      <211> 185
      <212> DNA
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      <220>
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      \langle 223 \rangle n = A,T,C or G
      <400> 26
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                                                                         60
agaatcttcn gggcgcgtca aaacaattgg gtgnattaag gacaanctcg gtcancagta
                                                                        120
taanctctct ttcncgngga ttantngnca taatcatnat tctgacnngt aggacattnc
                                                                        180
caacc
                                                                        185
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      <211> 270
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
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      <223> n = A,T,C or G
      <400> 27
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                                                                         60
atcacatctg gaaaccacca ntaccaccac cactacgcac ntcaccaaaa ctgtganagg
                                                                        120
gggcatttca gagacaanaa ttgaaaancg aatagtcntc acgggggnat gcanacattg
                                                                        180
accatgacca ggcgctggct caggcagnta aagaggccan agatcaacac cctgacatgt
                                                                        240
cngtgaccag agtggtggtc cttacanaga
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      <210> 28
      <211> 758
      <212> DNA
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      <220>
      <221> misc feature
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      <400> 28
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                                                                         60
aaagccgcaa ttacaatcaa gaggaaccta cttccctcct ggcaaagaaa cccaaggaag
                                                                        120
gcgagcggaa gatttacttg gcaattgaaa gtgccaatga actggctgtg cagaaagcaa
                                                                        180
aggcagaaat caccaggctc ataaaagaag agctgatccg gctgcaaaat tcataccaac
                                                                        240
caacaaataa aggaagatac aaagtcttat agacatccgg aaaaaagatt tttacctgtg
                                                                        300
ctggtctatg atgtatgtgg cagttgctgt ctgcagttta caatgtattg tnaatqaaqa
                                                                        360
ttttttaaat tctatcttgc tgatttttt taaatataan aaactggtac ttggtaaaga
                                                                        420
aatctgtccg taattncccc ccaatcagtc caactatatt taaagccacc tgttttcnaa
                                                                        480
ttttgatntc ctttaatgtt nactccaata tccatatttt aaatgtcccg gataatatcc
                                                                        540
caaaggttta aaaaatggaa atntttgaac ttcnnttgaa nanaataaat tcccatcctt
                                                                        600
```

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tangggntnt ccccttnccc gttcttccaa gaaatgtgac cttccccaaa aaagntnatc
                                                                        660
cctanctttt tgnttccccc ctgantttct gancccggac antnacgggt ttaaaaanttt
                                                                        720
                                                                        758
ttaaattttc caanncaaaa aaccntntnn ttttttna
      <210> 29
      <211> 577
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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      <223> n = A,T,C or G
      <400> 29
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gatngaggtg aaaacgatat tgatccntct ggggttttac ggtgtgcact gggtgctgca
                                                                        120
                                                                        180
cnnacttqtc aaggtttgnt acgtcctctg ggcatctgca aaaggccctg ctctctggag
tqttqtatgt agtgtaccaa aanagtattt atacatccca ccaatcaaaa cacagctttn
                                                                        240
ttacctcatg cgaactcatn caaaccaata gaatntcaac atgttctgta ccttanagtg
                                                                        300
                                                                        360
ctcacttact acctctgaac natactcacg ctgtnntttg tctcttnctt atctttttgc
ntcttqtaat taactctttg tttcccttca tcaaatgtaa tgtanatcgt gatctattaa
                                                                        420
aanaaaaatc anggttgcac ttgctacttt naanaaaccg antgtggaaa cattgggtct
                                                                        480
naattcacac aggatcngta naactgttgt ggatactgag aaacntttga atgttcctcc
                                                                        540
ccttattacc atcccgcaaa aaaacccctn tnntttt
                                                                        577
      <210> 30
      <211> 449
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      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(449)
      <223> n = A, T, C or G
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                                                                         60
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aggganagat gaaaaattat aacnaagcat aatatagcaa ggactaaccc ctatnccttn
                                                                        120
tgcataatga attaactaga aataactttg caaggagagc caaagctaan accnccgaaa
                                                                        180
ccagacgagc tacctangaa cagctaaaaag agcacacccg tctatgtagc anaatagtgg
                                                                        240
gaagatttat aggtagaggc gacaaaccta ccgagcctgg tgatagctgg ttgtccaaga
                                                                        300
tagaatetta gttcaacttt aaatttgeee acanaaceet ataaateeee ttgtaaattt
                                                                        360
aactgttagt ccaaagagga acagctcttt ggacactagg aaaaaacctt gtagagagag
                                                                        420
                                                                        449
tcataaaaaa aanccctntn qqqnnnngn
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       <211> 500
       <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
       <222> (1)...(500)
       <223> n = A, T, C \text{ or } G
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<400> 31
tentggacce nggteecenn gngancaaan aagaagggen ngnttneatn gaaaaneetg
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tgattntcgc cccggtncag gtgttnannt atggcccncn cncatctggt atacgccnaa
                                                                        120
acaatntant tttacaatnn gtnccccanc aaacaangtt cgtngnnttn actaggtagt
                                                                        180
taatcccncc ccatgttcaa ataaagggcc cgcgntncna ataaggaanc cnccccgant
                                                                        240
ggggtccccg aggccctctc cttcataaaa nncattcaac ttccctcccn ctannaaagn
                                                                        300
aattnttcna atttttnaaa cactccctgt ccanggggac tttnccccca ntanctgaaa
                                                                        360
aaatngcntg acgttcccct tcggcctaag ggcncaactt anttnncccc caanacccgn
                                                                        420
gggnnaggnn naaactcccc tngaagggaa cnactcgcnt aaaaanggaa taatcncccc
                                                                        480
cnaattattc cctncccqqq
                                                                        500
      <210> 32
      <211> 426
      <212> DNA
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      <220>
      <221> misc_feature
      <222> (1)...(426)
      \langle 223 \rangle n = A,T,C or G
      <400> 32
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ctccctggga ccttttcccc ttcctgttta anaanccagg gctgcctgga ggaagctttg
                                                                        120
tcagatctag tggaatgtga cctccctgga atatgtgccc aggggtttgt ctaagcagtt
                                                                        180
teaggetaig geetttacte catetggtee ceatecetet tatetetete atgtgtgget
                                                                        240
gcacctggac gcttggacca tagctgtcac agccccctgg ggaggaaccc actccttggc
                                                                        300
cathtcagcc tgtgcaatgc aaggctcttg tttgatctgt gtgctgacan aaagcccagc
                                                                        360
ttccttaaga acttttcatg tggaacactt tggttttgan aagaaaataa atcanaaacc
                                                                        420
attaaa
                                                                        426
      <210> 33
      <211> 375
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(375)
      \langle 223 \rangle n = A,T,C or G
      <400> 33
ngttgcacct attggccngc tggtctcgac tcctgacctc gttatctgcc tgcctcggcc
                                                                         60
tectaaagtg etgggattae aggagtgage cacagtgeet ggeetgteaa gaettetett
                                                                        120
aagttaactt cctgagaagt gatgtctaaa agtatctttg ctggtgtgag aactccagtt
                                                                        180
tccaacacat attatttccc tcaactattt ggaatatttt agaattttaa ttccaaagga
                                                                        240
ttagtttgaa tacaagtatg ccacataact cagttttcgc catcttncat ttcttaacag
                                                                        300
tgtaaattaa aagctaataa tcataataat aaagtgcatt taattatctt cgaaaaaaaa
                                                                        360
aaancccttt tgggg
                                                                        375
      <210> 34
      <211> 809
      <212> DNA
      <213> Homo sapien
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<220>
     <221> misc feature
      <222> (1)...(809)
     <223> n = A,T,C or G
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tcccaaagtg ctggaactac aggtgtgagc caccacgcct ggcagctttg tgttcttttc
                                                                        120
                                                                        180
tttctgtgat cttgccttag atcacacaga taaaacatga caggacctgg accttaacac
agtttggctc tcaatcctgt tctcataacc acnactgcct tcatttatct gtgtcatcct
                                                                        240
                                                                        300
caqacctqac acatagtagg tgctcagtca gtgttcacta agtaaatgat gaccaagaac
tetttgactg ggtccaaggt gettateeca atacttegee atggetacet eceteattee
                                                                        360
tragctgact tgctctctct agcctggctg ctcctatttt atttcctaaa catggaccca
                                                                        420
                                                                        480
tggcaataag tttaaancta acangttgat acggtaccca tccataattt aatnaattnt
ggggctcatg caaccncaaa aaccagaacc caaaactacc tgtncncaaa caacaatcat
                                                                        540
ttinggtngg gatccentne tngcttggne cetttttta aaatgteeat teeceeegga
                                                                        600
ctttaagaaa ttgaaggaat ncccggaaan tattgttanc gggccccctt nagngaaaaa
                                                                        660
ggtggcnctc cnnncggggg ccctccctgt ccctgaaatt tnaaaacccc cctcccnntt
                                                                        720
                                                                        780
taaanccctt aatcccggnt aacancnaaa naaaattcta gggcccaaac ccannggttt
                                                                        809
ggttttaaaa aaccntntat ttttttnat
      <210> 35
      <211> 192
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(192)
      <223> n = A, T, C \text{ or } G
      <400> 35
caccttattg ggatacagca gtgaattaag ctattaaaat aagataatga ttgcttttat
                                                                         60
accttcagta gagaaaagtc tttgcatata aagtaatgtt taaaaaaacat gtattgaaca
                                                                        120
cgacattgta tgaagcacaa taaagattct gaagccaaaa aaaaaaaccc caanggggnt
                                                                        180
                                                                        192
nnttttnaaa aa
      <210> 36
      <211> 368
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(368)
      \langle 223 \rangle n = A,T,C or G
      <400> 36
ctgctagtac caantattat ttaagantac ttttcactac tcctaaataa tgacacagat
                                                                         60
acgtttgtct tacacatttc actttattgt caagttatta gtatgtttat tttcaaaagt
                                                                        120
tattttttgc aatttctttt tattattccg tactttttaa atttacttca ttatcacgtc
                                                                        180
ttcctttatt ctttttaaat agtttttgct tttgttattt tgttttccct tttttactct
                                                                        240
tggtttgtaa tacctctttc cttatttgct cctttctcat ttgatctcaa tgttaatcca
                                                                        300
actqttttcc acatctgatt cactaaaatt ttagcccaaa aaaaaaancc cntttngggg
                                                                        360
```

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gngntttt
                                                                         368
      <210> 37
      <211> 219
      <212> DNA
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      <220>
      <221> misc_feature
      <222> (1)...(219)
      \langle 223 \rangle n = A,T,C or G
      <400> 37
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                                                                          60
tacnatecea tenaentgea ectatanene ttecaetaeg cacateacea aanetgtgaa
                                                                         120
agggggcntn tcnttagaca cacaattgca gaatngacnn cncancccgg gggannctcn
                                                                         180
angttcaccn tgnagcaggn gctggctcan gctnttata
                                                                         219
      <210> 38
      <211> 198
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(198)
      <223> n = A, T, C \text{ or } G
      <400> 38
tcgatacagg gncagatctg ggagccaggg cgttgctgat gagttgcaca gacgatcaca
                                                                          60
totgaaacca coagtaccac caccactacg cacatcacca aagcgotggo tonggoaatt
                                                                         120
aangaggcca aagagcanca ccctgacatg tengtgacen ttgtantggt centaangac
                                                                         180
acngacatcg cctccaca
                                                                         198
      <210> 39
      <211> 560
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(560)
      <223> n = A, T, C \text{ or } G
      <400> 39
tttnnatcng nacagctagt cctntaaant aatgacttca tagaaatggc attataattt
                                                                          60
ttaagttgat actctacagg tagctattga tataattagt tttaataaaa catgctgcaa
                                                                         120
ccatggtata caacaaaaat acatttcttt ggtgattgaa attaaggccg tatttacaat
                                                                         180
gacttaatat aagactgact tttatcctgc ttcataactt gtatggagaa ctcaccaaga
                                                                         240
aagaattcaa tactgtgaaa tatgcagcaa gaagattggt ctttacctag gctgtgtttc
                                                                         300
ctaagctctg agttttcagc accagtagat ttgtattaaa agaaaaaaa atggggcctt
                                                                         360
agcttctggc ttttaatttt gccagctaag gacataaaac aaaantaanc aancaaaanc
                                                                         420
aaatagccat ntgctatcag catcattatg taaaagaaaa tntattttag cccctaaaat
                                                                         480
taggaagaat gtaatctcag aataaaggtt gtcatttaag ttgaataaat atntagcttt
                                                                         540
cgaaaaaaa aanccccttt
                                                                         560
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<210> 40
      <211> 421
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1) ... (421)
      \langle 223 \rangle n = A,T,C or G
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tcgggataag atccaggcat gncttttaaa tctcagaggt agcagtaaac ttttcantnt
                                                                        120
tgcngttagc aagtgtgtgt ttgccaataa anccccatta tactaatgtg cctanttaat
                                                                        180
gttcagggaa natctgcttc cactgtgtnc cnaggggtgn catgaactnt gtgagnagcc
                                                                        240
concnnotgg agggatgaat gotgngttaa ctacngctat cacggatngt gtgntgtgaa
                                                                        300
naatacatcn acatnaatnt tanntgctct gnaanttccc ttnttatntg tcaagtaact
                                                                        360
                                                                        420
ntttgtaaaa ntnntnctcc caanttatta cngtgattac taatnnattn gtnccatgtt
                                                                        421
      <210> 41
      <211> 411
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(411)
      <223> n = A,T,C \text{ or } G
      <400> 41
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                                                                          60
aagatggact gggaaaaaca tcaactcctg aagttagaaa taagaatggt ttgtaaaatc
                                                                         120
cacagctata tcctgatgct ggatggtatt aatcttgtgt agtcttcaac tggttagtgt
                                                                         180
                                                                         240
gaaatagttc tgccacctct gacgcaccac tgccaatgct gtacgtactg catttgcccc
ttgagccagg tggatgttta ccgtgtgtta tataacttcc tggctccttc actgaacatg
                                                                         300
cctantccaa catttttcc cagtggagtc ncatcctggg atccagtgta taaatcccaa
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ttatcatgtc ttgtgcataa attcttccca aaagggatct ntaatttttt g
                                                                         411
      <210> 42
      <211> 408
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(408)
      <223> n = A, T, C \text{ or } G
      <400> 42
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                                                                          60
ccctgaatct cctgacaaat gcgaacagga actcctattc atcaggagcc aacttgataa
                                                                         120
                                                                         180
ctganaagat tectetetea tttateagee tttgattate tttttgtgte tettaetatt
tgcgcttagc gagaaaaata aagaggtttg aacaattaag aagtaacaaa gagctcatag
                                                                         240
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ttcacaaaga gcaantcaaa ggatgtctgg aatatttgaa catacaactg cctttggcat 300 gaggtggcct acatacattc tcaggggcag gataggctgg nanagctgat caagctgccg 360 ggaaagctga agcaaaggca gggttggntg gaaatcaaaa tntctctt 408 <210> 43 <211> 275 <212> DNA <213> Homo sapien <220> <221> misc_feature <222> (1)...(275) <223> n = A, T, C or G<400> 43 tecetaacte tetaagtaet teeettaeee acteagtgtg gtgatggeac eteeetgaat 60 ctcctgacaa atgcgaacag gaactcctat tcatcagagc caacttgata actgagaaga 120 ttcctctctc atttatcage ctttgattat ctttttgtgt ctcttactat ttgcgcttag 180 caagaaaaat aaagaggttt gaacaantaa gaagtancnn ggagctcnta gttcanaagn 240 agcaagtcaa aggatgtctg gangatttga agggt 275 <210> 44 <211> 246 <212> DNA <213> Homo sapien <220> <221> misc feature <222> (1)...(246) <223> n = A, T, C or G<400> 44 tttggtccca agcacatttc acaaangaga atttacacct agcacagctg gtgccangan 60 athtectang gacatggeca cetgggteca etecagegae agacecetga caagageagg 120 tctctggagg ctnantngca tggggcctan tntcntcaat cnaatgagcc ccnantgcta 180 ctgcgccccg ggggctccca cggcctgggc nnctttcntg caactgnaaa aggatagngg 240 tatttc 246 <210> 45 <211> 345 <212> DNA <213> Homo sapien <220> <221> misc feature <222> (1)...(345) <223> n = A, T, C or G<400> 45 tttggctccg tgggacgttg tantgtgcnc agacatttcc aagggaaatt ctaaacagtc 60 accetneect titgeattee eccaaatett aagtgtatae ataaaaceet gggtacatat 120 tgtngtggta atagaaggga attggnnaaa cngtacactt gttatatgga antnactgtg 180 gccacctaca aaagacaagt taacaaactg tentggagge tgtngntgee canccaggge 240 cgctgcnttt tgacaacatt cccaccctgg ccactcagca canttcatgg caggtcatgt 300 ctntncactg anachtttnt ganacttttt catatagcan aatcc 345

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<210> 46
     <211> 969
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(969)
      <223> n = A.T.C or G
      <400> 46
aattqcaqtt ctttcttgcc tttaacaaca ttagggcctt tagaatgagt acctggtgct
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gtocttocaa ctotgtgatt ototgattoo atootoattt ttoaccatoa otggtgtact
                                                                       120
                                                                       180
qqcaaqaacc antatgagat ttgaggaaaa atacttggat tactctttt taaaaaaaat
tatttagata taattcccat accatacaat taaccttttt atgtgtataa ttcagtattt
                                                                       240
ntagtatate cacaaagttg tgetaceate accaetatee gatteeagag ettgteatea
                                                                       300
tacaaaaaaa aaaaccccan agtnanttcc tttcaaaacn ctttnngttn ttcnttntnc
                                                                       360
centgtngen tetagnneng ngggntnnet tttgtenntn tenecetnen eteatentnn
                                                                       420
                                                                       480
enggtetetg etengngnnn egntntgnet tnnanteget getnntentg tatteeeege
netngtnnng tetgennegt agecagtggn ceteetgntn cennengntt etntntnegg
                                                                       540
cacannteca necanetgee atnagtnana nnatetetnt tenneanetg ntnncagnnt
                                                                       600
                                                                       660
tqtcntcntc tccqtnccnc cngcngctnn ctcnttncgc nctggnngnc antcgtacct
ggettttate eccetnteen netnttetng atggnntete ntetenacae etgnegttae
                                                                       720
gnntctcntn tnncnnnann cgttnctntn tnncttnccg ncngccatct nagctcannc
                                                                       780
                                                                       840
tgqngcgant cncgctctgn gtatcagtca tntanagann ngngnntgtt nccnncgcgn
nntgaganne conceenett egeatnacgt angtgnettt ntnnatetge tegtegtete
                                                                       900
nctcatatcc nccatgctgn catganactc cntantctnn cgcnnttctn ncgttccctc
                                                                       960
                                                                       969
tgcccttnn
      <210> 47
      <211> 361
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(361)
      <223> n = A, T, C or G
      <400> 47
                                                                        60
ggccactaag caggtcttac cnaatttaag aanattgaan tcctatcaag tatctcttct
gaccacaatg gtatgaaact agaaatcagt aacaggagga aaattggaag attcacaaat
                                                                       120
ntgtggaant taatcaacne atgagcaact antgagtena agancanate aaaagggann
                                                                       180
                                                                       240
tcaaaaactc tcttgaggtg gatgagaatg ganatacaac ataccngaac tcatgggatg
tatcacaagc ngtgctaagg gggaagttta agtnctagat gtctanatta ngaaagggaa
                                                                       300
agateteana tanacnacce agentinene etegaanaae tagaaaaaaet aagaaaaaaa
                                                                       360
                                                                       361
t
      <210> 48
      <211> 364
      <212> DNA
      <213> Homo sapien
      <220>
```

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```
<221> misc_feature
      <222> (1)...(364)
      <223> n = A, T, C \text{ or } G
      <400> 48
atgatgacca catntagatg gcacatngat gaggacttta atctttcctt aaanacaata
                                                                         60
atgtgttctt ttttctttta ntcacatgat ttctaagtan attttncatg caggacactt
                                                                        120
tttcaacctt gatgtacant gactgtgtaa aatttntctt tcagtggcaa cctctataat
                                                                        180
ctttannata tggtgagcat ctngtctgtt tagaanggga tatgacaata aatctatcag
                                                                        240
atggaaaatc ctgttacaaa gtataaaagc tttaqtaatt tactcaqtqt qqtqqtttta
                                                                        300
tcctttttgc tttttctccc ttggtctata atgaaattgt tacaqcaqtq caaaataaaa
                                                                        360
tcct
                                                                        364
      <210> 49
      <211> 703
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1) ... (703)
      <223> n = A, T, C or G
      <400> 49
atggggaatc aaacaatgtt aaaaggctan taatacttat aggttttatg attcaattta
                                                                         60
ctatgtgttt aaaattgttt tttgaaaaaa ttgagttatg tcnctaaaac tgagtctnta
                                                                        120
cagcicaaaa aigaagaaat achiatcicc gataagcata tiatqiqaat ticaacatch
                                                                        180
ctattgagaa aaggaatata aatttgaatg aaaatgaaac tctatctttc tatatcacat
                                                                        240
tgcataggtg taggctagtg agtactttga tgtaaattgc tgtatctttt gaggcntcna
                                                                        300
tttggcnata tagatcagaa ttttaaatcn gcatactttg tttgccagaa atctatcaqq
                                                                        360
accacttgta ntnattttgt tnaaaggaat atcnaacnct tggatgttca ncncagtatt
                                                                        420
gattgtttta naagaaggaa anggagaaag ggaggagaat ggaaqanana aanggagga
                                                                        480
ggaanattgg aaccnttgac atntgtgata gcatnggatt tgctnaacac nctatantat
                                                                        540
accectngea tggganaage atgeaenetn aaacaaggae nngttngatg gntetaennt
                                                                        600
ttgacntcag atnnaantaa atnaaaaaaa aaancccccn cctctttgnn ttcctntcnn
                                                                        660
cgnnnnannc ntctccccnc nncgnccnnc ncccgccacc ntn
                                                                        703
      <210> 50
      <211> 413
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(413)
      \langle 223 \rangle n = A,T,C or G
      <400> 50
tcttggctgg ttgagtattc aanaatcagg cacggagaag tggggtggat gcaaaccaac
                                                                         60
tgaccactgt ggcaccacca gcagtttcag ttttcatctt gantqtcnaq aqqaaatatc
                                                                        120
taatcttaca actcnttagg ggcctggctc agtggctcat accttgtntt cccancactt
                                                                        180
tgggangceg angenggent atcaccegca ngtcaggatt ttgagaccac cetggecaac
                                                                        240
ntggtgaaac cccatctcta ctantcaata caaancttag ctanqcgtga tqqcatqcac
                                                                        300
ctctaatccc acttacttgg gangctgagg cagcganaat cacttgtaac ccggaaggca
                                                                        360
nacgttgcat ntgagccaag atcgtgccac tgcactccat cctgggcttt cta
                                                                        413
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<210> 51
      <211> 252
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(252)
      <223> n = A,T,C or G
      <400> 51
gttacagaca aggnttntag aatatcttat gttttatgct ctgtaagttc aaagaagnta
                                                                        60
qcaqaaaaca taagcatact gaaaagagaa acagaagcta ttttttaaat acctatgtga
                                                                        120
aatctctcta tntgaaacaa aaaatacact ggatggatta gacactgcag aaggaaaatt
                                                                        180
                                                                        240
tqqtqaactt qagatcttat aaataaaaat tatccaaaat gaagtgtaga gtgaaaaaaa
                                                                        252
aaaancccct at
      <210> 52
      <211> 875
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(875)
      \langle 223 \rangle n = A,T,C or G
      <400> 52
agaaacgaga atgganattc aaatacgtcn gccgggcttg gtggattaga cctgtaaccc
                                                                         60
                                                                        120
naacactttg ggaggnctag gtgggcggat caccngaggt cnngagtacg ggaacancet
ggcaaaaacc ccntctttan tctgngaaaa cncaactcta ctaaaanaac tactcttaga
                                                                        180
                                                                        240
tnggcgtngn tgcgcctgcc tgttntccca gatacnnttt naggctgang tggggataan
tnctttaaca tgggaagtgg aagttgcact gatccaatgt ctccacactg cantccagcc
                                                                        300
tgggttangg aatgagaccc cncncacgga aaggacaata aaaanccccn nnggnnttnn
                                                                        360
tttttaangg cctcttgntc nttttcttnt antgcncgcc tncgcnnncn ttgntntgtc
                                                                        420
gantennntg ennttnttte ttennecten anectgette tnntenntte geenntnnae
                                                                        480
ngetteecc ntnetetage actinnitie thieghteen nnateteenn etintetnin
                                                                        540
cegetegegt nnncentnan etegnntent necetteett enengennen ntttegnena
                                                                        600
gateginegn etetatetae tietnieenn gnintanata ingaintiae atinigeten
                                                                        660
atnacccatn annnenteta tgtttatann ngtnnnneen tteaaennnn enttatgagn
                                                                        720
tettnactea getetnegtt gntntteena etanngttgn nentneatgt netgtenegt
                                                                        780
anchetethe tentenengt entgagaena atetetatht atnghttath cetgenthet
                                                                        840
                                                                        875
ganctncacc gngatctcgg cnntntcttc tcaag
      <210> 53
      <211> 182
      <212> DNA
      <213> Homo sapien
      <400> 53
ccagaagaag ggctacatat ggactcatgt tgggcctact cctgcaataa caattaagga
                                                                         60
atcagttgcc aaccatttgt agttcacaaa ttaaaactgg gtttccaggc ctggtgtgt
                                                                        120
                                                                        180
qqctcacqcc tgtagcccca gctattgcac cactgctctc caagctgggc aatggagtca
                                                                        182
ga
```

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```
<210> 54
      <211> 329
      <212> DNA
      <213> Homo sapien
      <400> 54
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                                                                     60
tagagagtca ggactagaag ttcagtctag ggatcaaata ataatagtag ctaatgttta
                                                                     120
aaggtaccta agatccgcca ggagacatac tcagtatagt tccgtggttt gccacatttc
                                                                     180
atcttatcca gtagcacagg tgaaatttgt cttatgtgta tactgaggaa aaacaagtcc
                                                                     240
ctctqatacc agcagccaat aaatgacaaa gctgggatag aaacttactt cattctaacc
                                                                     300
cgagagtccc tgttcttgca tggggcaca.
                                                                     329
      <210> 55
      <211> 312
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(312)
      <223> n = A, T, C \text{ or } G
      <400> 55
actuaacteg tittgagetat aggaatngge cattegnngt ggeteanace tgtaateeca
                                                                     60
gnatttnggg anacctcact aggatcacnt gaggtcagga gttcaagacc agcctgtcca
                                                                     120
acatggngaa accccatctc tantanaaaa tacagaaatt atccaggtgt ggtggctggc
                                                                     1.80
acctgtaatc ccagctactt gggaggccaa ggcatggaaa attgtctgaa cctqqqaaqt
                                                                     240
ggaggttgcg gtnanctgan atcatgccat tgctctccag cctcggccac anatcaagac
                                                                     300
cctatctcaa aa
                                                                     312
      <210> 56
      <211> 565
      <212> DNA
      <213> Homo sapien
      <400> 56
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                                                                     60
gaatttggcc ctcgaggcca agaattcggc acgaggggat ccaacgtcgc tccagctgct
                                                                     120
cttgacgact ccacagatac cccgaagcca tggcaagcaa gggcttgcag gacctgaagc
                                                                     180
aacaggtgga ggggaccgcc caggaagccg tgtcagcggc cggagcggca gctcagcaag
                                                                     240
tggtggacca ggccacagag gcggggcaga aagccatgga ccagctggcc aagaccaccc
                                                                     300
aggaaaccat cgacaagact gctaaccagg cctctgacac cttctctggg attgggaaaa
                                                                     360
aattcggcct cctgaaatga cagcagggag acttgggtcg gcctcctgaa atgacagcag
                                                                     420
ggagacttgg gtgacccccc ttccaggcgc catttagcac agcctggccc tgatctccqq
                                                                     480
540
aaaaaaaag atgcggccgc aagct
                                                                     565
      <210> 57
      <211> 798
      <212> DNA
      <213> Homo sapien
      <400> 57
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ggaacaagta gaagggaaga gaatgaggatg gagggataca gcttcccttct gaggaaggga actcacagca tcgcactccc gtgcagtgta aagagacaag gttgggcatta ctaacaaccc cctgcctgca tacatgtgaa catatgctta agacagagtg	gaagtetgea agaagtaatg agteettaag acetgateat etggteaagg caaaagetat eagetaetgg	cagctgtaaa aaagcacatg gcaaagggag ctgatcacac taaataggtt agaggacatg aatcttccag	ggttttatag tgaataaccc gcagtgctga ttgtgccaac gaacaatcaa caaattctac atttcagtgt	atgtettige ettecatece ageattggtg ttgatteata taacattate agteatteet tttaaaatea	60 120 180 240 300 360 420 480
gagctctgaa tacacaaaag g tgatactcag tgacaggagc a tctcagtaaa tcaagtccct a agaaatccag gaggcaatat g ccactagttt gcccttctat ctatagtgag tcgtatta	acagagetet tacetatgtt gtetttatte	aatgtccaca ctgacactga taatgaagtc	ggatgttgta ggctcttgga ctcatcttgc	gggtagggtc gctatgggtt actcagaggc	540 600 660 720 780 798
<210> 58 <211> 729 <212> DNA <213> Homo sapie <400> 58	n				
aagaatagac cgagataggg agaacgtgga ctccaacgtc gtgaaccatc accctaatca accctaaagg gagcccccga aggaagggaa gaaagcgaaa tgcgcgtaac caccacaccc attcaggctg cgcaactgtt gctggcgaaa gggggatgtg gtcacgacgt tgtaaaacga gggccctcta gatgcatgct ttgtaatacg actcactata ttgtaatggg tatggagaca ttatccgct <210> 59 <211> 730 <212> DNA	aaagggcgaa agttttttgg tttagagctt ggagcgggcg gccgcgctta gggaagggcg ctgcaaggcg cggccagtga cgagcggccg gggcttttt tatcatataa	aaaccgtcta ggtcgaggtg gacggggaaa ctagggcgct atgcgccgct atcqgtgcgg attaagttgg attgtaatac ccagtgtgat ttttttcggt	tcagggcgat ccgtaaagca gccggcgaac ggcaagtgta acagggcgcg gcctcttcgc gtaacgccag gactcactat ggatatctgc ttgaggggga	ggcccactac ctaaatcgga qtggcgagaa gcggtcacgc tccattcgcc tattacgcca ggttttccca agggcgaatt agaattcggc atgctggaga	60 120 180 240 300 360 420 480 540 600 660 720 729
<213> Homo sapie <400> 59 aagaatagac cgagataggg agaacgtgga ctccaacgtc gtgaaccatc accctaatca acctaaagg gagccccga aggaagggaa gaaagcgaaa tgcgcgtaac caccacaccc attcaggctg cgcaactgtt gctggcgaaa gggggatgtg gtcacgacgt tgtaaaacga gggcctcta gatgcatgct ttgtaatacg actcactata ttgtaatgg ttgtaatgg ttgtaatgg actcactata ttgtaatgg ttgtaatgg actcactata ttgtaatgg ttgtaatgg actcactata ttgtaatgg ttgtaatgg actcactata ttgtaatgg ttgtaatgg<	ttgagtgttg aaagggcgaa agttttttgg tttagagctt ggagcggcg gccycgctta gggaagggcg ctgcaaggcg cggccagtga cgagcggccg	aaaccgtcta ggtcgaggtg gacggggaaa ctagggcgct atgcgccgct atcggtgcgg attaagttgg attgtaatac ccagtgtgat	tcagggcgat ccgtaaagca gccggcgaac ggcaagtgta acagggcgcg gcctcttcgc gtaacgccag gactcactat ggatatctgc ttgagggga	ggcccactac ctaaatcgga gtggcgagaa gcggtcacgc tccattcgcc tattacgcca ggttttccca agggcgaatt agaattcggc atgctggaga	60 120 180 240 300 360 420 480 540 600 660 720 730

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<210> 60

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<211> 623
      <212> DNA
      <213> Homo sapien
      <400> 60
gactccaaga gaagactagg aagtagccct cgttctccag ggcacccaaa ataccagcct
                                                                      60
ttattgtctg catgatttta ggggatatgg ggagggaaca agtagaaggg aagagggaaa
                                                                     120
tggagagcat ccttatgact ttacaaaggg tggaaatgag gatggaggga tacagaagtc
                                                                     180
tgcacagctg taaaggtttt atagatgtct ttgccttccc ttctgaggaa gggaagaagt
                                                                     240
aatgaaagca catgtgaata accccttcca tcccattcac agcatcgcac tcccagtcct
                                                                     300
taaggcaaag ggaggcagtg ctgaagcatt ggtggtgcag tgtaaagaga caagacctga
                                                                     360
tcatctgatc acacttgtgc caacttgatt catattgggc attactaaca acccctgggc
                                                                     420
aaggtaaata ggttgaacaa tcaataacat tatccctgcc tgcatacatg tgaacaaaag
                                                                     480
ctatagagga catgcaaatt ctacagtcat teeteatatg etttagacag agtgcageta
                                                                     540
600
gccctatagt gagtcgtatt aca
                                                                     623
      <210> 61
      <211> 376
      <212> DNA
      <213> Homo sapien
      <400> 61
gcatgctcga gcggccgcca gtgtgatgga tatctgcaga attcggctta gcggataaca
                                                                      60
atttcacaca ggatccatga ctcagctant aaggetetgg cettggatee ctatgaggaa
                                                                     120
tattttacca caggitcagc agaaggtaac ataaaggitt ggagattgac aggccatggc
                                                                     180
ctaattcatt catttaaaag tgaacatgct aagcagtcca tatttcgaaa cattggggct
                                                                     240
ggagtcatgc agattgacat catccagggc aatcggctct tctcctgtgg tgcagatggc
                                                                     300
acgctgaaaa ccagggmttt gcccaatgct tttaacatcc ctaacagaat tcttgacatt
                                                                     360
ctataaagat tggggt
                                                                     376
      <210> 62
      <211> 539
      <212> DNA
      <213> Homo sapien
      <400> 62
atgactcatt gtttctctgc ctttccgtgt gttacaggtg ggctgatccc cctgcagcca
                                                                     60
gtttcccata agcaactgac ttccaactgg gaatgtctcg ggggataatg ggggtgggga
                                                                     120
tatggaagta tagagaaaac ataagaaaat actgggtgta tacacctttc tctctctgag
                                                                     180
tatgatgaca atgtgatagt cagtgtggca tctgcgactc cagcttgtgc ctggcatgta
                                                                     240
caccctaget ccagetteec etgggagaet gtgcatetee tggeteeact aacaccacet
                                                                     300
tottotgaco ttocagocta gagatgatga ototgocago otagatgggo totgggttgt
                                                                    360
ctccctattc ctgtttgctt tgtagatttc ccattatgct gtcaccaact ccccagccta
                                                                    420
agccctctct attttaaatt ctcaagtgga ttatgttcct gattagtccc tgactgatat
                                                                     480
accactetee teatgatete tgattagttt teetgttagg ttgttgeagt aaaaaaaaa
                                                                    539
      <210> 63
      <211> 304
      <212> DNA
      <213> Homo sapien
      <400> 63
ggcttagcgg ataacaattt cacacaggac gactccaagc tgggaaggaa aattcccttt
                                                                     60
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<210> 67

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tccaacctgt atcaattttt acaacttttt tcctgaaagc agtttagtcc atactttgca
                                                                         120
                                                                         180
ctgacatact ttttccttct gtgctaaggt aaggtatcca ccctcgatgc aatccacctt
gtgttttctt agggtggaat gtgatgttca gcagcaaact tgcaacagac tggccttctg
                                                                         240
                                                                         300
tttgttactt tcaaaaggcc cacatgatac aattagagaa ttcccaccgc acaaaaaaaa
                                                                         304
aaaq
      <210> 64
      <211> 226
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1) ... (226)
      \langle 223 \rangle n = A,T,C or G
      <400> 64
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                                                                          60
atqacactga tgattctcac cagtctgatg agtctcacca ttctgatgaa tctgatgaac
                                                                         120
tggtcactga ttttcccncg gacctgccng caaccgaagt nttcactcca gttgtccccc
                                                                         180
cagtagacac ntntgatggc cgaggtgatg gtgtggttta tggact
                                                                         226
      <210> 65
      <211> 225
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(225)
      <223> n = A,T,C or G
      <400> 65
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                                                                          60
                                                                         120
qaqqaaqatq atattganag aaggaaagaa ttgaaagcat cttgaagaaa aactcagatt
ggatntggga ttggtcaagt cggccggata atattccccc caaggagttc ctctttaaac
                                                                         180
                                                                         225
accegaageg caeggeeace eteageatga ggaacaegag egtea
      <210> 66
      <211> 240
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
       <222> (1)...(240)
       \langle 223 \rangle n = A,T,C or G
      <400> 66
                                                                          60
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gtacttgtgg aacaacatca aaaggtggca ggccatatac aaacagtacg acactgaccg
                                                                         120
atcagggacc atgtgcagta gtgaactccc angtgccttt gaggcagcan ggttccacct
                                                                         180
                                                                         24.0
qaatqaacan ctctataaca tgatcatccg acnctactca gatgaaagtg ggaacatgga
```

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<211> 504
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(504)
      \langle 223 \rangle n = A,T,C or G
      <400> 67
cacqaqqaqa gatngcatct gctatatatt ccacngatac atgtgagtna ctgatagaaa
                                                                      60
aaatcgcnnc ggngaacact gncaccggtn ccggcccccg gtactacagg qatctcntca
                                                                     120
gacttcaccg tntactacaa ngtaagcncc ctttaagaat gtcacggagt atgatgqqca
                                                                     180
ggatgcctgc ggctccaaca nctggaacnt ggtggacqtq qacctcccqc ccaacaaqqa
                                                                     240
cntggagccc ggcatcttac tacatgggct gaanccctqq actcaqtacq ccqtttacnt
                                                                     300
caaggctgtg accctcacca tggtggagaa cgaccatatc cgtggggcca agagtgagat
                                                                     360
cttgtncatt cgcnccantg cttcngttcc ttccnttccc ttgqacnttc tttcqqcatc
                                                                     420
aaactcctct tctcagttaa tcgtgaagtg gaaccctccc tctctgccca acggcnacct
                                                                     480
gagttactac tttgtgcnct ggca
                                                                     504
      <210> 68
      <211> 462
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(462)
      <223> n = A,T,C or G
      <400> 68
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                                                                      60
120
agagtgttgt ctctccccaa atttataaaa actaaaatgc atnccattcc tctgaaagca
                                                                     180
aaacaaattc ataattgagt gatattaaat anagaggttt tcggaagcag atctgtgaat
                                                                     240
atgaaataca tgtgcatatt tcattcccca ggcagacatt ttttaqaaat caatacatqc
                                                                     300
cccaatattg gaaagacttg ttcttccacg gtgactacag tacatgctga agcgtgccgt
                                                                     360
ttcagccctc atttaattca atttgtaagt agcgcagcag cctctgtggg ggaggatagg
                                                                     420
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                                                                     462
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      <211> 357
      <212> DNA
      <213> Homo sapien
      <400> 69
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ccaaagctaa aggatgatga ggttgctcag ctcaagaaaa gtqqaqatac cctqtqqqac
                                                                     120
atccagaagg acctaaaaga cctgtgacta gtgagctcta gqctqtagaa atttaaaaac
                                                                     180
tacaatgtat taactcgatc ctttagtttt catccatgta catggatcac agtttgcttt
                                                                     240
gatcttcttc aattgtgaat ttgggctcac agaatcaaag cctatgcttg qtttaatqct
                                                                     300
tgcaatctga gctcttgaac aaataaaatt aactattgta gtgtgaaaaa aaaaaaa
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      <210> 70
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<211> 226

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<212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(226)
      <223> n = A, T, C \text{ or } G
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atgacactga tgattctcac cagtctgatg agtctcacca ttctgatgaa tctgatgaac
                                                                        120
tggtcactga ttttcccncg gacctgccng caaccgaagt nttcactcca gttgtccccc
                                                                        180
                                                                        226
caqtagacac ntntgatggc cgaggtgatg gtgtggttta tggact
      <210> 71
      <211> 477
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(477)
      <223> n = A,T,C \text{ or } G
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cagtcgtcaa agctatttct ggagttcata ctgtcaggtt caaaaatgaa ctagaaagaa
                                                                         120
                                                                         180
atattacaat caagettgga tatgetaatg etaagattta taagettgat gacccaagtt
gccctcggcc agaatgttat agatcttgtg ggagcagtac acctgacgag tttcctacgg
                                                                         240
acattccagg gaccaaaggg aacttcagat tagtcagaca tgtttccttt gttgactgtc
                                                                         300
ctggccacna tattttgatg gctactatgc tgaacggtgc agcagtgatg gatgcagctc
                                                                         360
ttctgttgat agctggtaat gaatcttgcc ctcagcctca gacatcggaa acacctggct
                                                                         420
gctatagaag atcatgaaac tggaagccat attttgaatt ctacaaaata aaattga
                                                                         477
      <210> 72
      <211> 374
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc_feature
      <222> (1)...(374)
      <223> n = A, T, C \text{ or } G
      <400> 72
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                                                                          60
gtctcctcaa gattctctct atgggtctcg acacttaact gcaaagatgg catcgacccc
                                                                         120
gcacccacct ggagcgagag gcaccagcca actgcatggc atggatttat tggtcttatt
                                                                         180
ggatttgatt ggagctccaa acccaacgtt tcccaatttt tttccanact cagccaggtg
                                                                         240
gttcgaanga cttcaagcan ttgaacatga acttcatgaa ttgggtttgc tcaangatca
                                                                         300
ctctttggag gggcggtatt tccanaatta cagttatgga ggtgtgattc aggatgaccn
                                                                         360
                                                                         374
ttttccattt ccaa
      <210> 73
      <211> 597
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<212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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      \langle 223 \rangle n = A,T,C or G
      <400> 73
ccaagggatc tgtaaagaat atatacttga gtggtgtgtg ttatcagata aagcaccctg
                                                                         60
tatcacagac tggcaacaag aagatggtac cgtgcatcgc acctatttaa gagggaactt
                                                                         120
agcagagagc aaatgctatt tgataacagt tactccagta tatgctgatg gaccaggaag
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ccctgaatcc ataaaggcat accttaaaca agctccacct tccaaaggac ctactgttcg
                                                                        240
gacaaaaaa gtagggaaaa acgaagctgt cttanagtgg gaccaacttc ctgttgatgt
                                                                        300
tcanaatgga tttatcagaa attatactat attttatana accatcattg gaaatgaaac
                                                                        360
tgctgtgaat gtggattctt cccacacaga aatntacatt gtcctctttq actaqtqaca
                                                                        420
cattgtacat ggtacgaatg gcagcataca cagatgaagg tgggaaggat ggtccaaaat
                                                                        480
tcacttttac taccccaaan tttgctcaag gganaaattg aagccatant cgtgcctgtt
                                                                        540
tgcttancat tcctattgac aactcttctg ggaatgctgt tctgctttaa taagcga
                                                                        597
      <210> 74
      <211> 257
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(257)
      \langle 223 \rangle n = A,T,C or G
      <400> 74
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                                                                         60
atctgaaccn agatgtnaaa naagaaaatg ctttgaggct ttctaagcga tcctcctgtc
                                                                        120
taattincac cittgictgg atgcacactt cigaconogo tgccacaacc tgtggggtct
                                                                        180
gatgtgtccc ttgatgggtg cggccctcag ggactgcacc ctgacaagtg ttnaggcaan
                                                                        240
attcctttct tgtgccc
                                                                        257
      <210> 75
      <211> 330
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(330)
      <223> n = A, T, C \text{ or } G
      <400> 75
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                                                                         60
ctgtgtatgg aggtcgnagc cacaatacgc ggacgangat gtgaacacct acaatgccqc
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catchettae accatectea gecaagatee tgageteeet gachaaaata tgttenecat
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taacaggaac gcaggagtca tcggtgtggt cnccactggg ctggaccgaa agagtttccc
                                                                        240
tacgtgtacc ntggtggttc aagcngctga ccttcanggt gaggggttaa tcacnacagc
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ancngctgtg atcacagtca ctgntaccaa
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```

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<210> 76
      <211> 387
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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      \langle 223 \rangle n = A,T,C or G
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tatctccaag atgctattcg ttgaacccat cctggaggtt tccagcttgc cgacaaccaa
                                                                        120
ctcaacaacc aattcagcca ccaaaataac agctaatacc actgatgaac ccaccacaca
                                                                        180
acceaceaca qaqeecacea cecaacecac catecaacec acceaaceaa etacecaget
                                                                        240
                                                                        300
cccaacaqat tetectacee ageceactae tgggteette tgeecaggae etgttactet
                                                                        360
ctgctctgac ttgganantc attcaacana agccgtgttg ggggaagctt tggtaaattt
                                                                        387
ctccctgaag ctctaccacg ccttctc
      <210> 77
      <211> 339
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1) ... (339)
      <223> n = A,T,C or G
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                                                                        120
tgatctgcgc cctggtcctg gtgtccatnc tggccctcgg nancctggcc gaggcccana
                                                                        180
canagacqtq tncaqtqqcc ccccqtgaaa gacagaattg tggttttcct ggtgtcacac
cctcccantg tgcaaataag ggctgctgtt tcgacaacac cgttcgtggg gtcccctggt
                                                                        240
                                                                        300
gcttctatcc taatacente nacntecene canaaaagga ntgtgaattt tanacaette
tgcagggatc tgcctgcatc ctgacgcngt gccgtcccc
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      <210> 78
      <211> 385
      <212> DNA
      <213> Homo sapien
      <400> 78
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                                                                         60
                                                                        120
tqatttttqt attgaatatt gctgtctgtt acaaagtcag ttaaaggtac gttttaatat
ttaagttatt ctatcttgga gataaaatct gtatgtgcaa ttcaccggta ttaccagttt
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attatqtaaa caagagattt ggcatgacat gttctgtatg tttcagggaa aaatgtcttt
                                                                        240
                                                                        300
aatqcttttt caagaactaa cacagttatt cctatactgg attttaggtc tctgaagaac
tgctggtgtt taggaataag aatgtgcatg aagcctaaaa taccaagaaa gcttatactg
                                                                        360
                                                                        385
aatttaaqca aaaaaaaaaa acccc
      <210> 79
      <211> 307
      <212> DNA
      <213> Homo sapien
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<220>
      <221> misc_feature
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      <223> n = A,T,C or G
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tcctgacaaa gnagaaagtc atctactctc acttcacatg tgctacagat acagacaata
                                                                        120
ttcgctttgt gtttgctgct gtcaaagaca caattctaca gctaanccta agggaattca
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accttgtcta aaagctgctg cccactcctc ccctataaca gaagatgtga tttgcaaact
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ccttgtttta tttgnaagtg cttctgacat cnccagagcc agccccatgc caggaactaa
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ggatgtc
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      <210> 80
      <211> 528
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
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      \langle 223 \rangle n = A,T,C or G
      <400> 80
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tgccctgcgt gacgtacgtc agcaatatga aagtgtggct gccaagaacc tgcaggaggc
                                                                       120
agaagaatgg tacaaatcca agtttgctga cctctctgag gctgccaacc ggaacaatga
                                                                       180
cgccctgcgc caggcaaagc aggagtccac tgagtaccgg agacaggtgc agtccctcac
                                                                       240
ctgtgaagtg gatgccctta aaggaaccaa tgagtccctg gaacgccaga tgcgttgaaa
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tggaagagaa ctttgccgtt gaagctgcta actaccaaga cactattggc cgcctgcagg
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atgagattca gaatatgaag ganggaaatg gctcgtcacc ttcgtgaata ccaagacctg
                                                                       420
ctcaatgtta agatggccct tgacattgaa attgccacct acanggaact gctggangen
                                                                       480
aagaaaacca ggatttctct gcctcctccn aacttttcct cccctgaa
                                                                       528
      <210> 81
      <211> 369
      <212> DNA
      <213> Homo sapien
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cgaaatcata tggctgtagc attcgtgcta tccctggggg ttgcagcttt gtataagttt
                                                                       120
cgtgtggctg atcaaagaaa gaaggcatac gcagatttct acagaaacta cgatgtcatg
                                                                       180
aaagattttg aggagatgag gaaggctggt atctttcaga gtgtaaagta atcttggaat
                                                                       240
ataaagaatt tottoaggtt gaattaccta gaagtttgto actgacttgt gttootgaac
                                                                       300
tatgacacat gaatatgtgg gctaagaaat agttcctctt gataaataaa caattaacaa
                                                                       360
aaaaaaaa
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      <210> 82
      <211> 269
      <212> DNA
      <213> Homo sapien
      <220>
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<221> misc feature
      <222> (1)...(269)
      \langle 223 \rangle n = A,T,C or G
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                                                                        120
atggaagaag tagactaatc tetggetgag ggatgaetta eetgtteagt actetacaat
                                                                        180
tcctctgata atatatttc aaggatgttt ttctttattt ttgttaatat taaaangtct
                                                                         240
                                                                         269
qtntqqnatq acaactnctt taaggggaa
      <210> 83
      <211> 196
      <212> DNA
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      <221> misc_feature
      <222> (1)...(196)
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                                                                         60
                                                                         120
nnttgctaaa ccttcccagg tgtattttgg aggtacagtt gttggcnagc aagctatnaa
atctgaagat gaagtgggaa gttnaatana gtatgaatnc agggtaagaa actnaggtaa
                                                                         180
                                                                         196
acctcnaata tncctc
      <210> 84
      <211> 448
      <212> DNA
      <213> Homo sapien
      <220>
      <221> misc feature
      <222> (1)...(448)
      <223> n = A, T, C \text{ or } G
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tnngatttcc accatatcna ncntcnggaa tttaaccntc aggagnagct cttnntcaga
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cnccctggaa aaacgagccc cattgnancc anctttgana cataaaacct ggagaaattc
                                                                         180
tccaatacng aaggtatana gcggggcatc gttgacagca tcacgggtca aaggcttctg
                                                                         240
gaggeteagg cetgeaaagg tggeateate cacecaacea egggeeagaa cetgtenett
                                                                         300
caggacgcag tctcccnggg tgtgattgac caagacatgg ccaccaggct gaagcctgct
                                                                         360
cagaaagcct tcataggctt cgagggtgtg aagggaaaga agaagatgtc agcagcagag
                                                                         420
                                                                         448
gcagtgaaaa aaaaaaaacc cctatatt
       <210> 85
       <211> 169
       <212> DNA
       <213> Homo sapien
       <400> 85
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                                                                          60
qaqtattagg aaacatgagc agcatatggc ttttgatcag tttttcagtg gcagcatcca
                                                                         120
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🗜 atgaacaaga teetacaage tgtgeaggea aaacetagea ggaaaaaaa

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